

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



WO 99/26966

3 June 1999 (03.06.99)

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) (51) International Patent Classification 6: (11) International Publication Number: **A2** C07K 14/00 (43) International Publication Date: (21) International Application Number: PCT/US98/25296 (22) International Filing Date: 25 November 1998 (25.11.98) Anderson Street, San Francisco, CA 94110 (US). SHIAU, Andrew, K. [US/US]; 34 Hugo Street #3, San Francisco, CA 94122 (US). (30) Priority Data: US 26 November 1997 (26.11.97) 08/980,115 CA 94306-2155 (US) et al. (63) Related by Continuation (CON) or Continuation-in-Part (CIP) to Earlier Application 08/980,115 (CON) US Filed on 26 November 1997 (26.11.97)

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(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### **Published**

Without international search report and to be republished upon receipt of that report.

# (54) Title: NUCLEAR RECEPTOR LIGANDS AND LIGAND BINDING DOMAINS

# (57) Abstract

The present invention provides new methods, particularly computational methods, and compositions for the generation of nuclear receptor synthetic ligands based on the three dimensional structure of nuclear receptors, particularly the thyroid receptor (herein referred to as "TR"). Also provided are crystals, nuclear receptor synthetic ligands, and related methods.

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# NUCLEAR RECEPTOR LIGANDS AND LIGAND BINDING DOMAINS

### **ACKNOWLEDGMENTS**

This invention was supported in part by grants from the National Institutes of 5 Health grant number 1 R01 DK 43787 and 5 R01 DK 41842. The U.S. Government may have certain rights in this invention.

# CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit of the following provisional applications: 10 United States Ser. No. 60/008,540 and 60/008,543, filed December 13, 1995, and Ser. No. 60/008,606, filed December 14, 1995. This application claims the benefit of the following U.S. patent application: United States Ser. No. 08/764,870, filed December 13, 1996.

#### INTRODUCTION

#### Technical Field

This invention relates to computational methods for designing ligands that bind to nuclear receptors, crystals of nuclear receptors, synthetic ligands of nuclear receptors and methods of using synthetic ligands.

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#### **Background**

Nuclear receptors represent a superfamily of proteins that specifically bind a physiologically relevant small molecule, such as hormone or vitamin. As a result of a molecule binding to a nuclear receptor, the nuclear receptor changes the ability of a cell to transcribe DNA, i.e. nuclear receptors modulate the transcription of DNA, although they may have transcription independent actions. Unlike integral membrane receptors and membrane associated receptors, the nuclear receptors reside in either the cytoplasm or nucleus of eukaryotic cells. Thus, nuclear receptors comprise a class of intracellular, soluble ligand-regulated transcription factors.

Nuclear receptors include receptors for glucocorticoids (GRs), androgens (ARs), mineralocorticoids (MRs), progestins (PRs), estrogens (ERs), thyroid hormones (TRs), vitamin D (VDRs), retinoids (RARs and RXRs), peroxisomes (XPARs and PPARs) and icosanoids (IRs). The so called "orphan receptors" are also

part of the nuclear receptor superfamily, as they are structurally homologous to the classic nuclear receptors, such as steroid and thyroid receptors. To date, ligands have not been identified with orphan receptors but it is likely that small molecule ligands will be discovered in the near future for this class of transcription factors. Generally, nuclear receptors specifically bind physiologically relevant small molecules with high affinity and apparent Kd's are commonly in the 0.01 - 20 nM range, depending on the nuclear receptor/ligand pair.

Development of synthetic ligands that specifically bind to nuclear receptors has been largely guided by the trial and error method of drug design despite the importance of nuclear receptors in a myriad of physiological processes and medical conditions such as hypertension, inflammation, hormone dependent cancers (e.g. breast and prostate cancer), modulation of reproductive organ function, hyperthyroidism, hypercholesterolemia and obesity. Previously, new ligands specific for nuclear receptors were discovered in the absence of information on the three dimensional structure of a nuclear receptor with a bound ligand. Before the present invention, researchers were essentially discovering nuclear receptor ligands by probing in the dark and without the ability to visualize how the amino acids of a nuclear receptor held a ligand in its grasp.

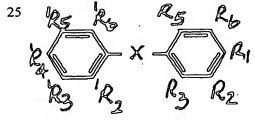
Consequently, it would be advantageous to devise methods and compositions 20 for reducing the time required to discover ligands to nuclear receptors, synthesize such compounds and administer such compounds to organisms to modulate physiological processes regulated by nuclear receptors.

# **SUMMARY OF THE INVENTION**

The present invention provides for crystals of nuclear receptor ligand binding domains with a ligand bound to the ligand binding domain (LBD). The crystals of the present invention provide excellent atomic resolution of the amino acids that interact with nuclear receptor ligand, especially thyroid receptor ligands. The three dimensional model of a nuclear receptor LBD with a ligand bound reveals a previously unknown structure for nuclear receptors and shows that the ligand is bound in a water inaccessible binding cavity of the ligand binding domain of the nuclear receptor.

The present invention also provides for computational methods using three dimensional models of nuclear receptors that are based on crystals of nuclear receptor LBDs. Generally, the computational method of designing a nuclear receptor ligand determines which amino acid or amino acids of a nuclear receptor LBD interact with a chemical moiety (at least one) of the ligand using a three dimensional model of a crystallized protein comprising a nuclear receptor LBD with a bound ligand, and selecting a chemical modification (at least one) of the chemical moiety to produce a second chemical moiety with a structure that either decreases or increases an interaction between the interacting amino acid and the second chemical moiety compared to the interaction between the interacting amino acid and the corresponding chemical moiety on the natural hormone.

Also provided is a method of modulating the activity of a nuclear receptor. The method can be *in vitro* or *in vivo*. The method comprises administering *in vitro* or *in vivo* a sufficient amount of a compound of the following formula:



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#### FORMULA I.

where the compound fits specially and preferentially into a nuclear hormone receptor LBD of interest. The method is exemplified by modulating the activity of a thyroid receptor (TR). For modulating TR activity, a compound of Formula I is employed

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PCT/US98/25296 WO 99/26966

that fits spacially and preferentially into a TR ligand binding domain (TR LBD), including compounds specific for a TR LBD isoform of interest. Of particular interest are the TR LBD isoforms  $\alpha$  (TR- $\alpha$ ) and  $\beta$  (TR- $\beta$ ). Additional compounds of interest include derivatives of Formula I, such as those compounds having the biphenyl (φ-X-5 φ) or single phenyl (φ-X or X-φ) nucleus of Formula I and its corresponding substituent groups described herein. Compounds that are interatively designed using structural information gleaned from these compounds and which modulate nuclear hormone receptor activity also are of interest.

The present invention also includes a method for identifying a compound 10 capable of selectively modulating the activity of a nuclear receptor. This aspect of the invention is exemplified by a method for identifying a compound capable of selectively modulating the activity of a TR isoform. The method comprises modeling test compounds that fit spacially and preferentially into a TR LBD isoform of interest using an atomic structural model of a TR LBD isoform bound to a test compound, 15 screening the test compounds in a biological assay for TR isoform activity characterized by binding of a test compound to a TR LBD isoform, and identifying a test compound that selectively modulates the activity of a TR isoform. compounds may be those of Formula I or derivatives thereof, including compounds having a biphenyl or single phenyl nucleus of Formula I.

Further included is a method for identifying agonist or antagonist ligands of a nuclear receptor using the atomic coordinates of a LBD in conjunction with a computerized modeling system. This aspect of the invention is exemplified by identifying a TR agonist or antagonist ligand by providing the atomic coordinates of a TR LBD to a computerized modeling system, modeling ligands which fit spacially 25 into the TR LBD, and identifying in a biological assay for TR activity a ligand which increases or decreases TR activity. The compounds can be those of Formula I or derivatives thereof, including compounds having a biphenyl or single phenyl nucleus of Formula I.

Also provided is a method of identifying a compound that selectively 30 modulates the activity of one type of nuclear receptor compared to other nuclear hormone receptors. The method is exemplified by modeling test compounds which fit spacially into a TR LBD using an atomic structural model of a TR LBD, selecting a compound comprising conformationally constrained structural features that interact

with conformationally constrained residues of a TR LBD, and identifying in a biological assay for TR activity a compound that selectively binds to a TR LBD compared to other nuclear receptors. The conformationally constrained features involved in receptor-selective ligand binding can be identified by comparing atomic models of receptor isoforms bound to the same and/or different ligands. The methods facilitate design and selection of compounds that have increased selectivity for a particular nuclear receptor. The compounds may be those of Formula I or derivatives thereof, including compounds having the biphenyl or single phenyl nucleus of Formula I.

Another aspect of the invention is a method for increasing the receptor 10 selectivity of a compound for a particular type of nuclear receptor. This involves the chemical modification of a substituent group of a compound of Formula I to generate compounds which have increased selectivity for one type of receptor. For example, chemical modification of a substituent group of the compound of Formula I can be 15 used to introduce additional constraints into a compound that modulates TR activity to increase its selectivity in vivo for TR-type receptors. Additional constraints also may The modified groups will preferably interact with a be added for stability. conformationally constrained structural feature of a TR LBD that is conserved among TR isoforms. A more preferred method comprises selecting compounds having 20 conformationally constrained groups that interact with conformationally constrained residues of a TR LBD conserved among TR isoforms. The compounds can be those of Formula I or derivatives thereof, including compounds having the biphenyl or single phenyl nucleus of Formula I.

The invention finds use in the selection and characterization of peptide,
25 peptidomimetic or synthetic compounds identified by the methods of the invention,
particularly new lead compounds useful in treating disorders related to nuclear
receptor-based deficiencies, including TR-related disorders. For TR-related disorders,
the compounds and methods of the invention can be used to modulate TR activity by
administering to a mammal in need thereof a sufficient amount of compound of
30 Formula I or derivative thereof that fits spacially and preferentially into a TR LBD.

# BRIEF DESCRIPTION OF THE DRAWINGS

- FIG. 1 is a diagram illustrating computational methods for designing ligands that interact with nuclear receptors of the nuclear receptor superfamily.
- FIG. 2 is a schematic representation of nuclear receptor structures, indicating 5 regions of homology within family members and functions of the various domains.
  - FIG. 3 shows the aligned amino acid sequences of the ligand binding domains of several members of the nuclear receptor superfamily.
- FIG. 4 is a ribbon drawing of the rat TR-α LBD with secondary structure elements labelled. The ligand (magenta) is depicted as a space-filling model. Alpha 10 helices and coil conformations are yellow, beta strands are blue.
  - FIG. 5 shows two cross-sections of a space-filling model of rat TR- $\alpha$  exposing the ligand (magenta) tightly packed within the receptor.
- FIG. 6 is a schematic of the ligand binding cavity. Residues which interact with the ligand appear approximately at the site of interaction. Hydrogen bonds are shown as dashed lines between the bonding partners; distances for each bond are listed. Non-bonded contacts are shown as radial spokes which face toward interacting atoms.
- FIG. 7 is the distribution of crystallographic temperature factors in the refined rat TR-I LBD. The distribution is represented as a color gradation ranging from less 20 than 15 (dark blue) to greater than 35 (yellow-green).
- FIG. 8 is a ribbon drawing of the rat TR-α LBD showing the c-terminal activation domain to ligand. Residues which comprise the c-terminal activation domain (Pro393-Phe405) are depicted as a stick representation. Hydrophobic residues, particularly Phe401 and Phe405 (blue) face inwards toward the ligand.

  25 Glu403 (red) projects outward into the solvent.
  - FIG. 9 is an electrostatic potential surface of the rat TR-α LBD, calculated using GRAPH. Negative electrostatic potential is red; positive electrostatic potential is blue. The c-terminal activation domain forms a largely hydrophobic (white). The Glu403 is presented as a singular patch of negative charge (red).
- FIG. 10 is a diagram comparing agonists and antagonists for several nuclear receptors.
  - FIG. 11 is the synthetic scheme for preparation of TS1, TS2, TS3, TS4 and TS5.

- FIG. 12 is the synthetic scheme for preparation of TS6 and TS7.
- FIG. 13 is the synthetic scheme for preparation of TS8.
- FIG. 14 is the synthetic scheme for preparation of TS10.
- FIG. 15 depicts the chemical structures of several TR ligands.
- 5 FIG. 16 is a graph illustrating competition assays in which T<sub>3</sub> and Triac compete with labeled T<sub>3</sub> for binding to human TR-α or human TR-β.
  - FIG. 17 depicts a Scatchard analysis of labelled  $T_3$  binding to  $TR-\alpha$  and  $TR-\beta$ .
- FIG. 18 is a chart showing the effect of TS-10 on the transcriptional regulation of the DR4-ALP reporter gene in the presence or absence of T3 as assayed 10 in TRAFI1 reporter cells.
  - FIG. 19 is a chart showing the effect of TS-10 on the transcriptional regulation of the DR4-ALP reporter gene in the presence or absence of T3 as assayed in TRAF91 reporter cells.
- FIG. 20 is a chart showing the effect of TS-10 on the transcriptional 15 regulation of the DR4-ALP reporter gene in the presence or absence of T3 as assayed in HepG2, a liver reporter cell line.
  - FIG. 21 is a partial ribbon drawing of TR-α LBD with T3 in the ligand binding cavity. Selected interacting amino acids are labelled, including Ile221, Ile222 and Ser260, Ala263, Ile299 and Leu 276.
- FIG. 22 is a partial ribbon drawing of TR-α LBD with T3 and Dimit superimposed in the ligand binding cavity. Interactions with Ile221, Ile222, Ala260, Ile 299 and Leu276 are labelled.
- FIG. 23 is a partial ribbon drawing of TR-α LBD with T3, illustrating the three Arginine residues (Arg228, Arg262 and Arg 266 (dark stick figures)) of the polar pocket, three water molecules HOH502, HOH503 and HOH504, with hydrogen bonds indicated by dotted lines.
  - FIG. 24 is a partial ribbon drawing of TR- $\alpha$  LBD with Triac, illustrating the three Arginine residues (dark stick figures) of the polar pocket, water molecules (HOH503, HOH504 and HOH600), with hydrogen bonds indicated by dotted lines.
- FIG. 25 is a partial ribbon drawing of the TR-α LBD with T3 and Triac superimposed in the ligand binding cavity. The drawing shows several interacting amino acid residues in the polar pocket that remain unchanged whether T3 or Triac occupies the ligand binding cavity: Arg262, Asn179, HOH503 and HOH504, and

Ser277. Both Arg228 and Arg 266 occupy two different positions, depending on whether T3 or Triac is bound.

- FIG. 26A and 26B are stereochemical representations of the TR- $\alpha$  LBD with Dimit bound.
- 5 FIG. 27 is a partial ribbon drawing of TR-β LBD with GC-1 in the ligand binding cavity. Amino acids Arg282, Arg316, Arg320, Asn 331 and His435 are labelled.
- FIG. 28 is a partial ribbon drawing of TR-β LBD with Triac in the ligand binding cavity. Amino acids Arg282, Arg316, Arg320, Asn331 and His435 are 10 labelled.
  - FIG. 29 is a partial ribbon drawing of TR-βLBD with GC-1 (Blue) overlayed with TR-α LBD with Dimit (Red) in the ligand binding cavities. Amino acids Arg228, Arg262, Arg266 and Ser277 (TR-α LBD), and Arg282, Arg316, Arg320 and Asn331 (TR-β LBD) are labelled.
- FIG. 30 is a partial ribbon drawing of TR-β LBD with Triac (Blue) overlayed with TR-α LBD with Triac (Red) in the ligand binding cavities. Amino acids Arg228, Arg262, Arg266, Ser277 and His381 (TR-α LBD), and Arg282, Arg316, Arg320 and His435 (TR-β LBD) are labelled.
- FIG. 31 is a graph showing competition curves comparing wildtype TR-α and 20 TR-β to a variant TR-β having a single amino acid substitution in the ligand binding domain.
  - FIG. 32 shows atomic numbering for thyronine-like ligands.
  - APPENDIX 1 is an appendix of references.
- **APPENDIX 2** is a chart of amino acids that interact with a TR ligand, for TR complexed with Dimit, Triac, IpBr2, T3 and GC-1.
  - APPENDIX 3 is a chart of atomic coordinates for the crystal of rat TR- $\alpha$  LBD complexed with Dimit.
  - APPENDIX 4 is a chart of atomic coordinates for the crystal of rat  $TR-\alpha$  LBD complexed with Triac.
- APPENDIX 5 is a chart of atomic coordinates for the crystal of rat TR-α LBD complexed with IpBr<sub>2</sub>.

APPENDIX 6 is a chart of atomic coordinates for the crystal of rat  $TR-\alpha$  LBD complexed with  $T_3$ .

APPENDIX 7 is a chart of atomic coordinates for the crystal of human TR- $\beta$  LBD complexed with Triac.

5 APPENDIX 8 is a chart of atomic coordinates for the crystal of human TR-β-LBD complexed with GC-1.

#### DETAILED DESCRIPTION OF THE INVENTION

#### INTRODUCTION

The present invention provides new methods, particularly computational methods, and compositions for the generation of nuclear receptor synthetic ligands based on the three dimensional structure of nuclear receptors, particularly the thyroid receptor (herein referred to as "TR"). Previously, the lack of three dimensional structural information about the ligand binding domain of a nuclear receptor thwarted the field of nuclear receptor drug discovery, especially the absence of three dimensional structural information relating to a nuclear receptor with a ligand bound.

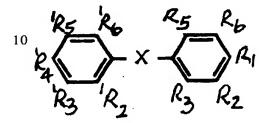
Described herein for the first time are crystals and three dimensional structural information from a nuclear receptor's ligand binding domain (LBD) with a ligand bound. The structure of the TR LBD complexed with 3,5,3'-triiodothyronine (T<sub>3</sub>), 20 3,5-dibromo-3'-isopropylthyronine (IpBr<sub>2</sub>), 3,5- dimethyl-3'-isopropylthyronine (Dimit), and 3,5,3'-triiodothyroacetic acid (Triac), 3,5-dimethyl-4-(4'-hydroxy-3'isopropylbenzyl)-phenoxy acetic acid (GC1) are exemplified. Such crystals offer superior resolution at the atomic level and the ability to visualize the coordination of nuclear receptor ligands by amino acids that comprise the LBD. The present 25 invention also provides computational methods for designing nuclear receptor synthetic ligands using such crystal and three dimensional structural information to generate synthetic ligands that modulate the conformational changes of a nuclear receptor's LBD. Such synthetic ligands can be designed using the computational methods described herein and shown, in part, in FIG. 1. These computational 30 methods are particularly useful in designing an antagonist or partial agonist to a nuclear receptor, wherein the antagonist or partial agonist has an extended moiety that prevents any one of a number of ligand-induced molecular events that alter the receptor's influence on the regulation of gene expression, such as preventing the

Page 12 of 447

WO 99/26966 PCT/US98/25296

normal coordination of the activation domain observed for a naturally occurring ligand or other ligands that mimic the naturally occurring ligand, such as an agonist. As described herein, synthetic ligands of nuclear receptors will be useful in modulating nuclear receptor activity in a variety of medical conditions.

Of particular interest is use of such ligands in a method of modulating TR activity in a mammal by administering to a mammal in need thereof a sufficient amount of a compound of Formula I,



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where the compound fits spatially and preferentially into a TR LBD. By "fits spacially" is intended that the three-dimensional structure of a compound is accommodated geometrically by a cavity or pocket of a TR LBD. By "TR LBD" is intended a structural segment or segments of thyroid hormone receptor polypeptide 20 chain folded in such a way so as to give the proper geometry and amino acid residue configuration for ligand binding. This is the physical arrangement of protein atoms in three-dimensional space forming a ligand binding pocket or cavity. By "fits spacially and preferentially" is intended that a compound possesses a three-dimensional structure and conformation for selectively interacting with a TR LBD. Compounds of 25 interest also include derivatives of Formula I. By "derivatives of Formula I" is intended compounds that comprise at least a single phenyl scaffold (\$\phi\$-X or X-\$\phi\$) of the biphenyl scaffold (φ-X-φ) of Formula I which comprise the corresponding substituents of Formula I described herein. Compounds that are interatively designed using structural information gleaned from these compounds and which 30 modulate nuclear hormone receptor activity also are of interest. Preferred compounds of Formula I and its derivatives that fit spacially and preferentially into a TR LBD comprise the following substituents:

(i) an R1-substituent comprising an anionic group that interacts with a side chain nitrogen atom of an arginine corresponding to a residue from the group Arg228, Arg262, and Arg266 of human TR-α, and Arg282, Arg316 and Arg320 of human TR-β, where the anionic group is about 1.7-4.0Å from the nitrogen atom;

- 5 (ii) an R2-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- (iii) an R3-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a serine, alanine and/or isoleucine corresponding to a residue from the group Ser260, Ala263 and Ile299 of human TR-α, and Ser314,
   10 Ala317 and Ile352 of human TR-β, where the hydrophobic or hydrophilic group is about 1.7-4.0Å from the side chain atom;
- (iv) an R5-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a phenylalanine and/or isoleucine corresponding to a residue from the group Phe218, Ile221 and Ile222 of human TR-α, and Phe272,
  15 Ile275 and Ile276 of human TR-β, where the hydrophobic or hydrophilic group is about 1.7-4.0Å from the side chain atom;
  - (v) an R6-substitutent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- (vi) an X-substituent comprising a hydrophobic or hydrophilic group that 20 interacts with a side chain atom of a leucine corresponding to a residue from the group Leu276 and Leu292 of human TR-α, and Leu 330 and Leu346 of human TR-β, where the hydrophobic or hydrophilic group is about 1.7-4.0Å from the side chain atom;
- (vii) an R2'-substituent comprising a hydrophobic or hydrophilic group that 25 fits spacially into the TR LBD;
- (viii) an R3'-substituent comprising a hydrophobic group that interacts with a side chain atom of a phenylalanine, glycine and/or methionine corresponding to a residue from the group Phe215, Gly290, and Met388 of human TR-α, and Phe269, Gly344, Met442 of human TR-β, where the hydrophobic group is about 1.7-4.0Å 30 from the side chain atom:
  - (ix) an R4'-substituent comprising an hydrogen bond donor or acceptor group that interacts with a side chain carbon or nitrogen atom of a histidine corresponding to residue His381 of human  $TR-\alpha$ , and His435 of human  $TR-\beta$ , where

the hydrogen bond donor or acceptor group is about 1.7-4.0Å from the side chain atom;

- (x) an R5'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- 5 (xi) and R6'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD; and

where the compound is other than thyronine (T3), triiodothyronine (T4) or other thyronine-like compounds previously known and used in a TR treatment method, such as those referenced in Appendix I.

10 Examples of such substituents include the following: where  $R_1$  is

-CH<sub>2</sub>CH[NH-tBOC]CO<sub>2</sub>H, or a carboxylate connected to the ring with a 0 to 3 carbon linker,

-PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CHNH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH[NHCOCHφ<sub>2</sub>]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH[NH-FMOC]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub> CH[NH-tBOC]PO<sub>3</sub>H<sub>2</sub>, or a phosphate or

20 phosphonate connected to the ring with a 0 to 3 carbon linker,

-SO<sub>3</sub>H, -CH<sub>2</sub>SO<sub>3</sub>H, -CH<sub>2</sub>CH<sub>2</sub>SO<sub>3</sub>H, -CH<sub>2</sub>CHNH<sub>2</sub>SO<sub>3</sub>H, -CH<sub>2</sub>CH[NHCOCH $\phi_2$ ]SO<sub>3</sub>H, -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>]SO<sub>3</sub>H, -CH<sub>2</sub>CH[NH-FMOC]SO<sub>3</sub>H, -CH<sub>2</sub> CH[NH-tBOC]SO<sub>3</sub>H, or a sulfate or sulfite connected to the ring with a 0 to 3 carbon linker, or acts as the functional

equivalent of CH<sub>2</sub>CH(NH<sub>2</sub>)CO<sub>2</sub>H of T3 in the molecular recognition domain when bound to a TR, wherein R<sub>1</sub> can be optionally substituted with an amine, where R<sub>2</sub> is

H, halogen, CF<sub>3</sub>, OH, NH<sub>2</sub>, SH, CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

30 where R<sub>3</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR, where  $R_5$  is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR, and  $R_3$  can be identical to  $R_5$ ,

#### where R6 is

5 -H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR, and R<sub>2</sub> can be identical to R<sub>6</sub>,

# where R2' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

where R<sub>3</sub>' is any hydrophobic group, including

halogen, -CF<sub>3</sub>, -SH, alkyl, aryl, 5- or 6-membered heterocyclie, cyano, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

#### 15 where R<sub>4</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, -SH, -CH<sub>3</sub>, -Et, or akyl, aryl or 5- or 6-membered heterocyclic aromatic attached through urea or carbamate linkages to O or N or S at the R<sub>4</sub>' position, or acts as the functional equivalent of OH in the molecular recognition domain when bound to a TR,

# where R<sub>5</sub>' is

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-H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups,

# where R<sub>6</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

where X is

O, S, SO<sub>2</sub>, NH, NR<sub>7</sub>, CH<sub>2</sub>, CHR<sub>7</sub>, CR<sub>7</sub>R<sub>7</sub>, wherein R<sub>7</sub> is alkyl, aryl or 5- or 6-membered heterocyclic aromatic,

and where the TR LBD ligand has an apparent Kd for binding TR LBD of 1 TM or 5 less.

Of particular interest are the class of compounds according to Formula I having the following substituents: where R<sub>1</sub> is carboxylate, phosphonate, phosphate or sulfite and is connected to the ring with a 0 to 3 carbon linker, R<sub>2</sub> is H, R<sub>3</sub> is -I, -Br, or -CH<sub>3</sub>, R<sub>5</sub> is -I, -Br, or -CH<sub>3</sub>, R<sub>6</sub> is H, R<sub>2</sub>' is H, R<sub>3</sub>' is -I, -Br, -CH<sub>3</sub>, -iPr, -phenyl, 10 benzyl, or 5- or 6-membered ring heterocycles, R<sub>4</sub>' is -OH, -NH<sub>2</sub>, and -SH, R<sub>5</sub>' is -H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally connected to the ring by a -CH<sub>2</sub>-, 15 aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups, and R<sub>6</sub>' is H.

The present invention also includes a method for identifying a compound capable of selectively modulating the activity of a TR isoform. By "modulating" is intended increasing or decreasing activity of a TR. By "TR isoform" is intended TR proteins encoded by subtype and variant TR genes. This includes TR-α and TR-β isoforms encoded by different genes (e.g., thra and thrb) and variants of the same genes (e.g., thrb1 and thrb2). The method comprises the steps of modeling test compounds that fit spacially and preferentially into a TR LBD isoform of interest using an atomic structural model of a TR LBD isoform bound to a test compound, screening the test compounds in a biological assay for TR isoform activity characterized by binding of a test compound to a TR LBD isoform, and identifying a test compound that selectively modulates the activity of a TR isoform. By "modeling" is intended quantitative and qualitative analysis of receptor-ligand structure/function based on three-dimensional structural information and receptor-ligand interaction models. This includes conventional numeric-based molecular

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WO 99/26966 PCT/US98/25296

dynamic and energy minimization models, interactive computer graphic models, modified molecular mechanics models, distance geometry and other structure-based constraint models. Modeling is preferably performed using a computer and may be further optimized using known methods.

5 For selectively modulating activity of a TR isoform, such as TR-α or TR-β, a sufficient amount of a compound that fits spatially and preferentially into TR LBD isoform is provided *in vitro* or *in vivo* to achieve the desired end result. TR-α isoform selectivity can be accomplished with a compound comprising an anionic group that interacts with an oxygen or carbon of a serine residue corresponding to Ser277 of 10 human TR-α, where the anionic group is about 1.7-4.0Å from the side chain atom. TR-β isoform selectivity can be accomplished with a compound comprising an anionic group that interacts with the side chain nitrogen of an asparagine corresponding to Asn331 of human TR-β, where the anionic group is about 1.7-4.0Å from the side chain nitrogen atom.

The present invention further includes a method for identifying a TR agonist or antagonist ligand by providing the atomic coordinates of a TR LBD to a computerized modeling system, modeling ligands which fit spacially into the TR LBD, and identifying in a biological assay for TR activity a ligand which increases or decreases the activity of the TR.

The invention also involves a method for increasing receptor selectivity of a compound of Formula I or derivatives thereof for a TR-type receptor versus other nuclear receptors by selecting a compound that interacts with conformationally constrained residues of a TR LBD that are conserved among TR isoforms. "Conformationally constrained" is intended to refer to the three-dimensional structure of a chemical or moiety thereof having certain rotations about its bonds fixed by various local geometric and physical-chemical constraints. In designing and selecting compounds having increased specificity for TRs compared to other nuclear receptors, the following methods of the invention can be used. One method involves comparing atomic models of a first TR LBD isoform bound to a compound with a second TR LBD isoform bound to the same compound, identifying atoms of the TR LBD and compounds which interact, and designing or selecting a compound that interacts with TR LBD residues comprising a conformationally constrained structural feature that is conserved between the TR LBD isoforms. Another method relates to comparing a

first TR LBD complexed with a first compound to a second TR LBD complexed with a second compound having one or more different substituents compared to the first compound, identifying atoms of the TR LBD and compounds which interact, and designing or selecting compounds that interact with TR LBD residues comprising a 5 conformationally constrained structural feature that is conserved between the TR LBD isoforms. The methods also facilitate identification of structural and conformationally constrained interactions that are conserved between compounds that bind to a TR LBD. The methods are exemplified by comparing atomic models of a first TR LBD isoform complexed with a first compound of Formula I to a second TR LBD isoform 10 complexed with the first compound, or a second compound of Formula I having different substituents than the first compound. For example, a TR-α LBD bound to a natural hormone such as T3 is compared to a TR-\$\beta\$ LBD bound to an organic thyronine-like compound such as GC-1. Conserved contacts are identified which are made between atoms of the different compounds and atoms of the TR LBDs, and the 15 fiducial and adjustable components identified. Compounds selective for TR are identified in a biological assay for TR activity that assays for selective binding to a TR and/or TR LBD compared to other nuclear receptors. Conventional assays for TR and other nuclear receptors may be conducted in parallel or serially, including those assays described herein. Automatable methods are preferred. The methods facilitate 20 design and selection of compounds comprising cyclic carbon and substituent atoms that interact with a constrained side chain and/or main chain atom of a TR LBD residue.

In another aspect of the invention, the methods described herein are useful for selecting peptides, peptidomimetics or synthetic molecules that modulate TR activity.

25 Methods of the invention also find use in characterizing structure/function relationships of natural and synthetic TR-ligands. Molecules of particular interest are new thyronine-like compounds other than T3, T4 and other thyronine-like compounds previously known and used for treating TR-related disorders. New compounds of the invention include those which bind to a TR LBD isoform with greater affinity than T3 or T4 and those which exhibit isoform-specific binding affinity.

#### APPLICABILITY TO NUCLEAR RECEPTORS

The present invention, particularly the computational methods, can be used to design drugs for a variety of nuclear receptors, such as receptors for glucocorticoids (GRs), androgens (ARs), mineralocorticoids (MRs), progestins (PRs), estrogens 5 (ERs), thyroid hormones (TRs), vitamin D (VDRs), retinoid (RARs and RXRs), icosanoid (IRs), and peroxisomes (XPARS and peroxisomal proliferators (PPAP)). The present invention can also be applied to the "orphan receptors," as they are structurally homologous in terms of modular domains and primary structure to classic nuclear receptors, such as steroid and thyroid receptors. The amino acid homologies 10 of orphan receptors with other nuclear receptors ranges from very low (<15%) to in the range of 35% when compared to rat RARI and human TR-β receptors, for example. In addition, as is revealed by the X-ray crystallographic structure of the TR and structural analysis disclosed herein, the overall folding of liganded superfamily members is likely to be similar. Although ligands have not been identified with 15 orphan receptors, once such ligands are identified one skilled in the art will be able to apply the present invention to the design and use of such ligands, as their overall structural modular motif will be similar to other nuclear receptors described herein.

### Modular Functional Domains Of Nuclear receptors

- The present invention will usually be applicable to all nuclear receptors, as discussed herein, in part, to the patterns of nuclear receptor activation, structure and modulation that have emerged as a consequence of determining the three dimensional structures of nuclear receptors with different ligands bound, notably the three dimensional structures or crystallized protein structure of the ligand binding domains 25 for TR-α and TR-β. Proteins of the nuclear receptor superfamily display substantial regions of amino acid homology, as described herein and known in the art see FIG. 2. Members of this family display an overall structural motif of three modular domains (which is similar to the TR three modular domain motif):
  - 1) a variable amino-terminal domain;
- 30 2) a highly conserved DNA-binding domain (DBD); and
  - 3) a less conserved carboxyl-terminal LBD.

The modularity of this superfamily permits different domains of each protein to separately accomplish different functions, although the domains can influence each

other. The separate function of a domain is usually preserved when a particular domain is isolated from the remainder of the protein. Using conventional protein chemistry techniques a modular domain can sometimes be separated from the parent protein. Using conventional molecular biology techniques each domain can usually be separately expressed with its original function intact or chimerics of two different nuclear receptors can be constructed, wherein the chimerics retain the properties of the individual functional domains of the respective nuclear receptors from which the chimerics were generated.

FIG. 2 provides a schematic representation of family member structures,
10 indicating regions of homology within family members and functions of the various domains.

#### Amino Terminal Domain

The amino terminal domain is the least conserved of the three domains and varies markedly in size among nuclear receptor superfamily members. For example, this domain contains 24 amino acids in the VDR and 603 amino acids in the MR. This domain is involved in transcriptional activation and in some cases its uniqueness may dictate selective receptor-DNA binding and activation of target genes by specific receptor isoforms. This domain can display synergistic and antagonistic interactions with the domains of the LBD. For example, studies with mutated and/or deleted receptors show positive cooperativity of the amino and carboxy terminal domains. In some cases, deletion of either of these domains will abolish the receptor's transcriptional activation functions.

#### 25 DNA-Binding Domain

The DBD is the most conserved structure in the nuclear receptor superfamily. It usually contains about 70 amino acids that fold into two zinc finger motifs, wherein a zinc ion coordinates four cysteines. DBDs contain two perpendicularly oriented I-helixes that extend from the base of the first and second zinc fingers. The two zinc 30 fingers function in concert along with non-zinc finger residues to direct nuclear receptors to specific target sites on DNA and to align receptor homodimer or heterodimer interfaces. Various amino acids in DBD influence spacing between two half-sites (usually comprised of six nucleotides) for receptor dimer binding. For

example, GR subfamily and ER homodimers bind to half-sites spaced by three nucleotides and oriented as palindromes. The optimal spacings facilitate cooperative interactions between DBDs, and D box residues are part of the dimerization interface. Other regions of the DBD facilitate DNA-protein and protein-protein interactions required for RXR homodimerization and heterodimerization on direct repeat elements.

The LBD may influence the DNA binding of the DBD, and the influence can also be regulated by ligand binding. For example, TR ligand binding influences the degree to which a TR binds to DNA as a monomer or dimer. Such dimerization also depends on the spacing and orientation of the DNA half sites. The receptors also can interact with other proteins and function to regulate gene expression.

The nuclear receptor superfamily has been subdivided into two subfamilies: 1) GR (GR, AR, MR and PR) and 2) TR (TR, VDR, RAR, RXR, and most orphan receptors) on the basis of DBD structures, interactions with heat shock proteins (hsp), and ability to form heterodimers. GR subgroup members are tightly bound by hsp in the absence of ligand, dimerize following ligand binding and dissociation of hsp, and show homology in the DNA half sites to which they bind. These half sites also tend to be arranged as palindromes. TR subgroup members tend to be bound to DNA or other chromatin molecules when unliganded, can bind to DNA as monomers and dimers, but tend to form heterodimers, and bind DNA elements with a variety of orientations and spacings of the half sites, and also show homology with respect to the nucleotide sequences of the half sites. By this classification, ER does not belong to either subfamily, since it resembles the GR subfamily in hsp interactions, and the TR subfamily in nuclear localization and DNA-binding properties.

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# Ligand Binding Domain

The LBD is the second most highly conserved domain in these receptors. Whereas integrity of several different LBD sub-domains is important for ligand binding, truncated molecules containing only the LBD retain normal ligand-binding activity. This domain also participates in other functions, including dimerization, nuclear translocation and transcriptional activation, as described herein. Importantly, this domain binds the ligand and undergoes ligand-induced conformational changes as detailed herein.

Most members of the superfamily, including orphan receptors, possess at least two transcription activation subdomains, one of which is constitutive and resides in the amino terminal domain (AF-1), and the other of which (AF-2 (also referenced as TAU 4)) resides in the ligand-binding domain whose activity is regulated by binding of an agonist ligand. The function of AF-2 requires an activation domain (also called transactivation domain) that is highly conserved among the receptor superfamily (approximately amino acids 1005 to 1022). Most LBDs contain an activation domain. Some mutations in this domain abolish AF-2 function, but leave ligand binding and other functions unaffected. Ligand binding allows the activation domain to serve as an interaction site for essential co-activator proteins that function to stimulate (or in some cases, inhibit) transcription.

For example, Shibata, H., et al. (Recent Progress in Hormone Res. 52:141-164 (1997)) has reviewed the role of co-activators and co-repressors in steroid/thyroid hormone receptor systems. Steroid receptor co-activator-one (SRC-1) appears to be a 15 general co-activator for all AF-2 domain containing receptors tested. SRC-1 enhances transactivation of steroid hormone-dependent target genes. Other putative coactivators have been reported, including the SRC-1 related proteins, TIF-2 and GRIP-1, and other putative unrelated co-activators such as ARA-70, Trip 1, RIP-140, and TIF-1. In addition another co-activator CREB-binding protein (CBP) has been shown 20 to enhance receptor-dependent target gene transcription. CBP and SRC-1 interact and synergistically enhance trancriptional activation by the ER and PR. A ternary complex of CBP, SRC-1, and liganded receptors-may form to increase the rate of hormone-responsive gene transcription. Co-repressors, such as SMRT and N-CoR, for TR and RAR, have been identified that also contribute to the silencing function of 25 unliganded TR. The unliganded TR and RAR have been shown to inhibit basal promoter activity; this silencing of target gene transcription by unliganded receptors is mediated by these co-repressors. The collective data suggests that upon binding of agonist, the receptor changes its conformation in the ligand-binding domain that enables recruitment of co-activators, which allows the receptor to interact with the 30 basal transcriptional machinery more efficiently and to activate transcription. In contrast, binding of antagonists induces a different conformational change in the receptor. Although some antagonist-bound receptors can dimerize and bind to their cognate DNA elements, they fail to dislodge the associated co-repressors, which results in a nonproductive interaction with the basal transcriptional machinery.

Similarly, the TR and RAR associate with co-repressors in the absence of ligand, thereby resulting in a negative interaction with the transcriptional machinery that silences target gene expression. In the case of mixed agonist/antagonists, such as 4-hydroxytamoxifen, activation of gene transcription may depend on the relative ratio of co-activators and co-repressors in the cell or cell-specific factors that determine the relative agonistic or antagonistic potential of different compounds. These co-activators and co-repressors appear to act as an accelerator and/or a brake that modulates transcriptional regulation of hormone-responsive target gene expression.

The carboxy-terminal activation subdomain, as described herein is in close three dimensional proximity in the LBD to the ligand, so as to allow for ligands bound to the LBD to coordinate (or interact) with amino acid(s) in the activation subdomain. As described herein, the LBD of a nuclear receptor can be expressed, crystallized, its three dimensional structure determined with a ligand bound (either using crystal data from the same receptor or a different receptor or a combination thereof), and computational methods used to design ligands to its LBD, including ligands that contain an extension moiety that coordinates the activation domain of the nuclear receptor.

Once a computationally designed ligand (CDL) is synthesized as described herein and known in the art, it can be tested using assays to establish its activity as an 20 agonist, partial agonist or antagonist, and affinity, as described herein. After such testing, the CDLs can be further refined by generating LBD crystals with a CDL bound to the LBD. The structure of the CDL can then be further refined using the chemical modification methods described herein for three dimensional models to improve the activity or affinity of the CDL and make second generation CDLs with 25 improved properties, such as that of a super agonist or antagonist described herein. Agonist and antagonist ligands also can be selected that modulate nuclear receptor responsive gene transcription through altering the interaction of co-activators and corepressors with their cognate nuclear hormone receptor. For example, CDL agonists can be selected that block or dissociate the co-repressor from interaction with the 30 receptor, and/or which promote binding or association of the co-activator. CDL antagonists can be selected that block co-activator interaction and/or promote corepressor interaction with the target receptor. Selection can be done in binding assays that screen for CDLs having the desired agonist or antagonist properties. Suitable assays for such screening are described herein and in Shibata, H., et al. (Recent Prog.

Horm. Res. 52:141-164 (1997)); Tagami, T., et al. (Mol. Cell Biol. 17(5):2642-2648 (1997)); Zhu, XG., et al. (J. Biol. Chem. 272(14):9048-9054 (1997)); Lin, B.C., et al. (Mol. Cell Biol. 17(10):6131-6138 (1997)); Kakizawa, T., et al. (J. Biol. Chem. 272(38):23799-23804 (1997)); and Chang, K. H., et al. (Proc. Natl. Acad. Sci. USA 94(17):9040-9045 (1997)), which references are incorporated herein in their entirety by reference.

#### **NUCLEAR RECEPTOR ISOFORMS**

The present invention also is applicable to generating new synthetic ligands to distinguish nuclear receptor isoforms. As described herein, CDLs can be generated that distinguish between binding isoforms, thereby allowing the generation of either tissue specific or function specific synthetic ligands. For instance, GR subfamily members have usually one receptor encoded by a single gene, although are exceptions. For example, there are two PR isoforms, A and B, translated from the same mRNA by alternate initiation from different AUG codons. There are two GR forms, one of which does not bind ligand. This method is especially applicable to the TR subfamily which usually has several receptors that are encoded by at least two (TR: α, β) or three (RAR, RXR, and PPAR: α, β, γ) genes or have alternate RNA splicing and such an example for TR is described herein.

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# **NUCLEAR RECEPTOR CRYSTALS**

The invention provides for crystals made from nuclear receptor ligand binding domains with the ligand bound to the receptor. As exemplified in the Examples, TRs are crystallized with a ligand bound to it. Crystals are made from purified nuclear receptor LBDs that are usually expressed by a cell culture, such as *E. coli*. Preferably, different crystals (co-crystals) for the same nuclear receptor are separately made using different ligands, such as a naturally occurring ligand and at least one bromo- or iodosubstituted synthetic ligand that acts as an analog or antagonist of the naturally occurring ligand. Such bromo- and iodo- substitutions act as heavy atom substitutions in nuclear receptor ligands and crystals of nuclear receptor proteins. This method has the advantage for phasing of the crystal in that it bypasses the need for obtaining traditional heavy metal derivatives. After the three dimensional structure is determined for the nuclear receptor LBD with its ligand bound, the three dimensional

structure can be used in computational methods to design a synthetic ligand for the nuclear receptor and further activity structure relationships can be determined through routine testing using the assays described herein and known in the art.

# 5 Expression and Purification of other Nuclear Receptor LBD Structures

High level expression of nuclear receptor LBDs can be obtained by the techniques described herein as well as others described in the literature. High level expression in E. coli of ligand binding domains of TR and other nuclear receptors, including members of the steroid/thyroid receptor superfamily, such as the receptors 10 ER, AR, MR, PR, RAR, RXR and VDR can also be achieved. Yeast and other eukaryotic expression systems can be used with nuclear receptors that bind heat shock proteins as these nuclear receptors are generally more difficult to express in bacteria, with the exception of ER, which can be expressed in bacteria. Representative nuclear receptors or their ligand binding domains have been cloned and sequenced: human 15 RAR-α, human RAR-γ, human RXR-α, human RXR-β, human PPAR-α, human PPAR-β, human PPAR-γ, human VDR, human ER (as described in Seielstad et al., Molecular Endocrinology, vol 9:647-658 (1995), incorporated herein by reference), human GR, human PR, human MR, and human AR. The ligand binding domain of each of these nuclear receptors has been identified and is shown in FIG. 3. Using the 20 information in FIG. 3 in conjunction with the methods described herein and known in the art, one of ordinary skill in the art could express and purify LBDs of any of the nuclear receptors, including those illustrated in FIG. 3, bind it to an appropriate ligand, and crystallize the nuclear receptor's LBD with a bound ligand.

FIG. 3 is an alignment of several members of the steroid/thyroid hormone 25 receptor superfamily that indicates the amino acids to be included in a suitable expression vector.

Extracts of expressing cells are a suitable source of receptor for purification and preparation of crystals of the chosen receptor. To obtain such expression, a vector is constructed in a manner similar to that employed for expression of the rat TR 30 alpha (Apriletti et al. Protein Expression and Purification, 6:363-370 (1995), herein incorporated by reference). The nucleotides encoding the amino acids encompassing the ligand binding domain of the receptor to be expressed, for example the estrogen receptor ligand binding domain (hER-LBD) (corresponding to R at position 725 to L

at position 1025 as standardly aligned as shown in the FIG. 3), are inserted into an expression vector such as the one employed by Apriletti et al (1995). For the purposes of obtaining material that will yield good crystals it is preferable to include at least the amino acids corresponding to human TR-β positions 725 to 1025.

5 Stretches of adjacent amino acid sequences may be included if more structural information is desired. Thus, an expression vector for the human estrogen receptor can be made by inserting nucleotides encoding amino acids from position 700 to the c-terminus at position 1071. Such a vector gives high yield of receptor in E. coli that can bind hormone (Seielstad et al. Molecular Endocrinology 9:647-658 (1995)).

10 However, the c-terminal region beyond position 1025 is subject to variable proteolysis and can advantageously be excluded from the construct, this technique of avoiding variable proteolysis can also be applied to other nuclear receptors.

# TR-α And TR-β As Examples of Nuclear receptor LBD Structure and Function 15 TR Expression, Purification And Crystallization

As an example of nuclear receptor structure of the ligand binding domain the α- and β- isoforms of TR are crystallized from proteins expressed from expression constructs, preferably constructs that can be expressed in E. coli. Other expression systems, such as yeast or other eukaryotic expression systems can be used. For the 20 TR, the LBD can be expressed without any portion of the DBD or amino-terminal domain. Portions of the DBD or amino-terminus can be included if further structural information with amino acids adjacent the LBD is desired. Generally, for the TR the LBD used for crystals will be less than 300 amino acids in length. Preferably, the TR LBD will be at least 150 amino acids in length, more preferably at least 200 amino acids in length, and most preferably at least 250 amino acids in length. For example the LBD used for crystallization can comprise amino acids spanning from Met 122 to Val 410 of the rat TR-α, Glu 202 to Asp 461 of the human TR-β.

Typically TR LBDs are purified to homogeneity for crystallization. Purity of TR LBDs is measured with sodium dodecyl sulfate polyacrylamide gel 30 electrophoresis (SDS-PAGE), mass spectrometry (MS) and hydrophobic high performance liquid chromatography (HPLC). The purified TR for crystallization should be at least 97.5 % pure or 97.5%, preferably at least 99.0% pure or 99.0% pure, more preferably at least 99.5% pure.

Initially purification of the unliganded receptor can be obtained by conventional techniques, such as hydrophobic interaction chromatography (HPLC), ion exchange chromatography (HPLC), and heparin affinity chromatography.

To achieve higher purification for improved crystals of nuclear receptors, 5 especially the TR subfamily and TR, it will be desirable to ligand shift purify the nuclear receptor using a column that separates the receptor according to charge, such as an ion exchange or hydrophobic interaction column, and then bind the eluted receptor with a ligand, especially an agonist. The ligand induces a change in the receptor's surface charge such that when re-chromatographed on the same column, 10 the receptor then elutes at the position of the liganded receptor are removed by the original column run with the unliganded receptor. Usually saturating concentrations of ligand are used in the column and the protein can be preincubated with the ligand prior to passing it over the column. The structural studies detailed herein indicate the general applicability of this technique for obtaining super-pure nuclear receptor LBDs for crystallization.

More recently developed methods involve engineering a "tag" such as with histidine placed on the end of the protein, such as on the amino terminus, and then using a nickle chelation column for purification, Janknecht R., *Proc. Natl. Acad. Sci. USA*, 88:8972-8976 (1991) incorporated by reference.

20 To determine the three dimensional structure of a TR LBD, or a LBD from another member of the nuclear receptor superfamily, it is desirable to co-crystalize the LBD with a corresponding LBD ligand. In the case of TR LBD, it is preferable to separately co-crystalize it with ligands such as T3, IpBr and Dimit that differ in the heavy atoms which they contain. Other TR ligands such as those encompassed by 25 Formula 1 described herein and known in the prior art, can also be used for the generation of co-crystals of TR LBD and TR ligands. Of the compounds encompassed by Formula 1 it is generally desirable to use at least one ligand that has at least one bromo- or iodo- substitution at the R3, R5, R3' or R5' position, preferably such compounds will be have at least two such substitutions and more preferably at 30 least 3 such substitutions. As described herein, such substitutions are advantageously used as heavy atoms to help solve the phase problem for the three dimensional structure of the TR LBD and can be used as a generalized method of phasing using a halogen (e.g. I or Br) substituted ligand, especially for nuclear receptors.

Typically purified LBD, such as TR LBD, is equilibrated at a saturating concentration of ligand at a temperature that preserves the integrity of the protein. Ligand equilibration can be established between 2 and 37°C, although the receptor tends to be more stable in the 2-20°C range.

Preferably crystals are made with the hanging drop methods detailed herein. Regulated temperature control is desirable to improve crystal stability and quality. Temperatures between 4 and 25°C are generally used and it is often preferable to test crystallization over a range of temperatures. In the case of TR it is preferable to use crystallization temperatures from 18 to 25°C, more preferably 20 to 23°C, and most preferably 22°C.

Complexes of the TR-\alpha LBD with a variety of agonists, including T<sub>3</sub>, IpBr<sub>2</sub>, Dimit, and Triac, are prepared with by methods described herein. For example, cocrystals of the rTR-α LBD, with ligand prebound, are prepared by vapor diffusion at ambient temperature from 15% 2-methyl-2,4-pentanediol (MPD). The crystals are 15 radiation sensitive, and require freezing to measure complete diffraction data. On a rotating anode X-ray source, the crystals diffract to ~3Å; synchrotron radiation extends the resolution limit significantly, to as high as 2.0Å for T<sub>3</sub> cocrystals. The composition of the thyroid hormone, combined with the ability to prepare and cocrystallize the receptor complexed with a variety of analogs, permitted the unusual 20 phasing strategy. This phasing strategy can be applied to the ligands of the nuclear receptors described therein by generating I and Br substitutions of such ligands. In this strategy, cocrystals of the TR LBD containing four hormone analogs that differ at the 3,5, and 3' positions (T<sub>3</sub>, IpBr<sub>2</sub>, Dimit, and Triac) provided isomorphous derivatives. For this set of analogs, the halogen substituents (2Br and 3I atoms) 25 function as heavy atoms, while the Dimit cocrystal (3 alkyl groups) acts as the parent. The initial 2.5Å multiple isomorphous replacement/anomalous scattering/density modified electron density map allowed the LBD to be traced from skeletons created in the molecular graphics program O5 (Jones, T.A. et al., ACTA Cryst, 47:110-119 (1991), incorporated by reference herein). A model of the LBD was built in four 30 fragments, Arg157-Gly184, Trp186-Gly197, Ser199-Pro205, and Val210-Phe405, and refined in XPLOR using positional refinement and simulated annealing protocols. Missing residues were built with the aid of difference density. The final model was refined to  $R_{cryst} = 21.8\%$  and  $R_{free} = 24.4\%$  for data from 15.0 to 2.2Å, see Table 6.

The human TR- $\beta$  LBD model was resolved by molecular replacement of the TR- $\alpha$  LBD coordinates. The structure is based on E202 to D461 with a his-tag at the N-terminus. The final model was refined to R<sub>cryst</sub> = 25.3% and R<sub>free</sub> = 28.9% for data from 30.0 to 2.4Å+, see Table 7.

This phasing strategy can be applied to the ligands of the nuclear receptors described herein by generating I and Br substitutions of such ligands.

# THREE DIMENSIONAL STRUCTURE OF TR LBD Architecture of TR LBD

As an example of the three dimensional structure of a nuclear receptor, the folding of the TR-α LBD is shown in FIG. 4. The TR-α LBD consists of a single structural domain packed in three layers, composed of twelve α-helices, H1-12, and four short β-strands, S1-4, forming a mixed β-sheet. The buried hormone and three antiparallel α-helices, H5-6, H9, and H10, form the central layer of the domain, as 15 shown in FIG. 4. H1, H2, H3 and S1 form one face of the LBD, with the opposite face formed by H7, H8, H11, and H12. The first 35 amino acids of the N-terminus (Met122-Gln156) are not visible in the electron density maps. The three dimensional structure of the heterodimeric RXR:TR DNA-binding domains bound to DNA, amino acids Met 122 - Gln151 of the TR DBD make extensive contacts with the minor 20 groove of the DNA8. The five disordered amino acids (Arg152-Gln156), which reside between the last visible residue of the TR DBD and the first visible residue of the LBD likely represent the effective "hinge" linking the LBD and the DBD in the intact receptor.

The predominantly helical composition and the layered arrangement of 25 secondary structure is identical to that of the unliganded hRXRa, confirming the existence of a common nuclear receptor fold between two nuclear receptors.

The TR LBD is visible beginning at Arg157, and continues in an extended coil conformation to the start of H1. A turn of α-helix, H2, covers the hormone binding cavity, immediately followed by short β-strand, S1, which forms the edge of the 30 mixed β-sheet, parallel to S4, the outermost of the three antiparallel strands. The chain is mostly irregular until H3 begins, antiparallel to H1. H3 bends at Ile221 and Ile222, residues which contact the ligand. The chain turns almost 90° at the end of H3 to form an incomplete α-helix, H4. The first buried core helix, H5-6, follows, its axis

altered by a kink near the ligand at Gly 253. The helix is composed of mostly hydrophobic sidechains interrupted by two striking exceptions: Arg262 is solvent inaccessible and interacts with the ligand carboxylate (1-substituent), and Glu256 meets Arg329 from H9 and Arg375 from H11 in a polar invagination. 5 terminates in a short β-strand, S2, of the four strand mixed sheet. S3 and S4 are joined through a left-handed turn, and further linked by a salt bridge between Lys284 and Asp272. Following S4, H7 and H8 form an L, stabilized by a salt bridge between Lys268 and Asp277. The turn between H7 and H8 adopts an unusual conformation, a result of interaction with ligand and its glycine rich sequence. H9 is the second core 10 helix, antiparallel to the neighboring H5-6. Again, two buried polar sidechains are found, Glu315 and Gln320. Glu315 forms a buried salt bridge with His358 and Arg356. The oxygen of Gln320 forms a hydrogen bond with the buried sidechain of His 175. The chain then switches back again to form H10, also antiparallel to H9. H11 extends diagonally across the full length of the molecule. Immediately after H11, the 15 chain forms a type II turn, at approximately 90° to H11. The chain then turns again to form H 12, which packs loosely against H3 and H11 as part of the hormone or ligand binding cavity. The final five amino acids at the C-terminus, Glu406 -Val410, are disordered. The architecture of the TR- $\beta$  LBD is identical to that of the TR- $\alpha$  LBD, with two significant differences. An additional helix is present at the N-terminus 20 (residues Glu202-I1e208), which is part of the DBD, and packs antiparallel to H10. Following the helix is a two residue turn (Gly209-His210) continuing into an extended coil to he start of H1, as seen in the TR-α LBD. A further difference occurs in the irregular conformation adopted between H2 and H3. In the TR-\alpha LBD, residue Gly197-Asp211 form a loop that packs against the receptor, contacting helices H7, 25 H8, H11, and the loop between H11 and H12. In the TR-β LBD, only the ends of the loop are ordered, with the stretch Ala253-Lys263 disordered. In addition to these residues, the residues of the His-tag at the N-terminus, and the final residue at the Cterminus, Asp461, are disordered.

# 30 TR LBD's Ligand Binding Cavity As An Example Of A Nuclear Receptor's Buried Ligand Cavity

The three dimensional structure of the TR LBD leads to the startling finding that ligand binding cavity of the LBD is solvent inaccessible when a T3 or its isostere

is bound to the LBD. This surprising result leads to a new model of nuclear receptor three dimensional structure and function, as further described herein, particularly in the sections elucidating the computational methods of ligand design and the application of such methods to designing nuclear receptor synthetic ligands that 5 contain extended positions that prevent normal activation of the activation domain.

Dimit, the ligand bound to the receptor, is an isostere of T<sub>3</sub> and a thyroid hormone agonist. Therefore the binding of Dimit should reflect that of T<sub>3</sub>, and the Dimit-bound receptor is expected to be the active conformation of TR. The ligand is buried within the receptor, providing the hydrophobic core for a subdomain of the 10 protein, as shown in FIG. 5 a and b. H5-6 and H9 comprise the hydrophobic core for the rest of the receptor.

An extensive binding cavity is constructed from several structural elements. The cavity is enclosed from above by H5-6 (Met 256- Arg266), from below by H7 and H8 and the intervening loop (Leu287- Ile299), and along the sides by H2 (185- 187), by the turn between S3 and S4 (Leu276-Ser277), by H3 (Phe215-Arg228), by H11 (His381-Met388) and by H12 (Phe401-Phe405). The volume of the cavity defined by these elements, calculated by GRASP (Columbia University, USA) (600 Å3), is essentially the volume of the hormone (530 Å). The change in volume can be exploited for ligand design as described herein. The remaining volume is occupied by water molecules surrounding the amino-propionic acid substituent. FIG. 6 depicts various contacts (or interactions) between TR's LBD and the ligand.

The planes of the inner and outer (prime ring) rings of the ligand are rotated from planarity about 60° with respect to each other, adopting the 3'-distal conformation (in which the 3' substituent of the outer ring projects down and away 25 from the inner ring). The amino-propionic acid and the outer phenolic ring assume the transoid conformation, each on opposite sides of the inner ring. The torsion angle χ<sub>1</sub> for the amino- propionic acid is 300°.

The amino-propionic acid substituent is packed loosely in a polar pocket formed by side chains from H2, H4 and S3. The carboxylate group forms direct 30 hydrogen bonds with the guanidium group of Arg228 and the amino N of Ser277. In addition, Arg262, Arg266 and Asn179 interact with the carboxylate through water-mediated hydrogen bonds. The three arginine residues create a significantly positive local electrostatic potential, which may stabilize the negative charge of the

carboxylate. No hydrogen bond is formed by the amino nitrogen. The interactions of the amino-propionic acid substituent are consistent with the fact that Triac, which lacks the amino nitrogen, has a binding affinity equal to that of T<sub>3</sub>, indicating that the amino nitrogen and longer aliphatic chain of T<sub>3</sub> do not contribute greatly to binding 5 affinity.

The biphenyl ether, in contrast, is found buried within the hydrophobic core. The inner ring packs in a hydrophobic pocket formed by H3, H5-6, and S3. Pockets for the 3- and 5-methyl substituents are not completely filled, as expected since the van der waals radius of methyl substituent for Dimit is smaller than the iodine substituent provided by the thyroid hormone T<sub>3</sub>. Such pockets are typically 25 to 100 cubic angstroms (although smaller pocket for substitutes are contemplated in the 40 to 80 cubic angstrom range) and could be filled more tightly with better fitting chemical substitutions, as described herein.

The outer ring packed tightly in a pocket formed by H3, H5-6, H7, H8, H11 and H12, and the loop between H7 and H8. The ether oxygen is found in a hydrophobic environment defined by Phe218, Leu287, Leu276, and Leu292. The absence of a hydrogen bond to the ether oxygen is consistent with its role in establishing the correct stereochemistry of the phenyl rings, as suggested by potent binding of hormone analogs with structurally similar linkages possessing reduced or negligible hydrogen bonding capability. The 3'-isopropyl substituent contacts Gly290 and 291. The presence of glycine at this position in the pocket can explain the observed relationship between activity and the size of 3'-substituents. Activity is highest for 3'-isopropyl, and decreases with added bulk. The only hydrogen bond in the hydrophobic cavity is formed between the phenolic hydroxyl and His381 Ne2.

The conformation of His381 is stabilized by packing contacts provided by Phe405, and Met256.

The presence of a 5' substituent larger than hydrogen affects the binding affinity for hormone. The more abundant thyroid hormone, 3,5,3',5'-tetraiodo-L-thyronine (T<sub>4</sub>), contains an iodine at this position, and binds the receptor with 2% of 30 the affinity of T<sub>3</sub>. The structure suggests that discrimination against T<sub>4</sub> is accomplished through the combination of steric conflict by Met256 and possibly the constraints imposed by the geometry of the hydrogen bond from His381 to the phenolic hydroxyl. The 5' position is a preferred location for introducing a chemical

modification of C-H at the 5' of T3 or and TR agonist, as described herein, that produces an extension from the prime ring and results in the creation of an antagonist or partial agonist.

Deletion and antibody competition studies suggest the involvement of residues

Pro162 to Val202 in ligand binding. The region does not directly contact hormone in
the bound structure, although H2 packs against residues forming the polar pocket that
interacts with the amino-propionic acid group. One role for H2, then, is to stabilize
these residues in the bound state, H2, with β-strands S3 and S4, might also represent a
prevalent entry point for ligand, since the amino-propionic acid of the ligand is
oriented toward this region. Studies of receptor binding to T3 affinity matrices
demonstrate that only a linkage to the amino-propionic acid is tolerated, suggesting
that steric hindrance present in other linkages prevent binding. Furthermore, the
crystallographic temperature factors suggest the coil and β-strand region is most
flexible part of the domain FIG. 7. Participation of this region, part of the hinge
domain between the DBD and LBD, in binding hormone may provide structural
means for ligand binding to influence DNA binding, since parts of the Hinge domain
contact DNA.

# TR LBD Transcriptional Activation Helix As An Example Of A Nuclear Receptor 20 Activation Domain

In addition to the startling finding that the ligand binding cavity is solvent inaccessible when loaded with a ligand, the activation helix of TR LBD presents a surface to the ligand cavity for interaction between at least one amino acid and the bound ligand. The C-terminal 17 amino acids of the TR, referred to as the activation helix or AF-2 (an example of an LBD activation domain), are implicated in mediating hormone-dependent transcriptional activation. Although, mutations of key residues within the domain decrease ligand-dependent activation it was unclear until the present invention whether such mutations directly affected ligand coordination. Although some mutations of this domain have been noted to reduce or abolish ligand binding, other mutations in more distant sites of the LBD have a similar effect.

Activation domains among nuclear receptors display an analogous three dimensional relationship to the binding cavity, which is a region of the LBD that binds the molecular recognition domain of a ligand, i.e. the activation domain

presents a portion of itself to the binding cavity (but necessarily the molecular recognition domain of the ligand). Many nuclear receptors are expected to have such domains, including the retinoid receptors, RAR and RXR, the glucocorticoid receptor GR, and the estrogen receptor ER. Based upon the TR's sequence, the domain is proposed to adopt an amphipathic helical structure. β-sheet or mixed secondary structures, could be present as activation domains in less related nuclear receptors.

Within the activation domain, the highly conserved motif ΦΦΧΕΦΦ, where Φ represents a hydrophobic residue, is proposed to mediate interactions between the receptors and transcriptional coactivators. Several proteins have been identified which bind the TR in a hormone-dependent fashion. One of these, Trip1, is related to a putative yeast coactivator Sug1, and also interacts with both the C-terminal activation domain and a subset of the basal transcriptional machinery, suggesting a role in transactivation by the TR. Other proteins, such as RIP140, SRC1, (Onate, S.A. et. al., Science 270:1354-1357 (1995)) and TF-1 (see also Ledouarim, B., et. al., 15 EMBO J. 14:2020-2033 (1995)), and GRIP-1 (Heery, E., et al., Nature 387:733-736 (1997)) also interact with other nuclear receptors in a ligand dependent manner through the C-terminal domain. Binding of these proteins can be modulated using the TR ligands described herein especially those TR ligands with extensions that sterically hinder the interaction between the highly conserved motif and other 20 proteins.

The C-terminal activation domain of the TR forms an amphipathic helix, H12, which nestles loosely against the receptor to form part of the hormone binding cavity. The helix packs with the hydrophobic residues facing inward towards the hormone binding cavity, and the charged residues, including the highly-conserved glutamate, extending into the solvent, as shown in FIG. 8. The activation helix of TR LBD presents Phe 401 to the ligand binding cavity and permits direct coordination with the hormone i.e. such amino acids interact with the ligand forming a van der waals contact with the plane of the outer phenyl ring. Phe 405 also interacts with His 381, perhaps stabilizing its hydrogen bonding conformation, i.e. a favorable hydrogen bond interaction. Participation of Phe 401 and Phe 405 in binding hormone explains how mutation of these residues decreases hormone binding affinity. Furthermore, the impact of these mutations on activation likely derives from a role in stabilizing the domain in the bound structure through increased hydrogen bond interaction of dipole

interactions. Glu 403 extends into the solvent, emphasizing its critical role in transactivation. In its observed conformation, presented on the surface as an ordered residue, against a background of predominantly hydrophobic surface, Glu 403 is available to interact with activator proteins described herein, as shown in FIG. 9. The other charged residues, Glu 405 and Asp 406 are disordered, as the helix frays at Phe 405.

Two other sequences in the TR, τ2 and τ3, activate transcription when expressed as fusion proteins with a DNA-binding domain. The sequences, discovered in the TRB, correspond to TR-α residues Pro158-Ile168 in H1 (τ2), and Gly290-Leu3 10 19 in H8 and H9 (τ3). Unlike the C-terminal activation domain, τ2 and τ3 do not appear to represent modular structural units in the rat TR-I LBD, nor present a surface for protein-protein interactions: the critical aspartate/glutamate residues of τ3 are located on two separate helices, and do not form a single surface; the charged residues of τ2 are engaged in ion pair interactions with residues of the LBD. Thus, τ2 and τ3 may not function as activation domains in the context of the entire receptor.

#### Computational Methods For Designing A Nuclear Receptor LBD LIGAND

The elucidation of the three dimensional structure of a nuclear receptor ligand binding domain provides an important and useful approach for designing ligands to nuclear receptors using the computational methods described herein. By inspecting the FIGURES it can be determined that the nuclear receptor ligand is bound in a water inaccessible binding cavity in the LBD and that chemical moieties can be added to selected positions on the ligand. Such chemical modifications, usually extensions, can fill up the binding cavity represented in the FIGURES for a tighter fit (or less water) or can be used to disrupt or make contacts with amino acids not in contact with the ligand before the chemical modification was introduced or represented in a figure of the three dimensional model of the LBD. Ligands that interact with nuclear superfamily members can act as agonists, antagonists and partial agonists based on what ligand-induced conformational changes take place.

Agonists induce changes in receptors that place them in an active conformation that allows them to influence transcription, either positively or negatively. There may be several different ligand-induced changes in the receptor's conformation.

Antagonists, bind to receptors, but fail to induce conformational changes that alter the receptor's transcriptional regulatory properties or physiologically relevant conformations. Binding of an antagonist can also block the binding and therefore the actions of an agonist.

Partial agonists bind to receptors and induce only part of the changes in the receptors that are induced by agonists. The differences can be qualitative or quantitative. Thus, a partial agonist may induce some of the conformation changes induced by agonists, but not others, or it may only induce certain changes to a limited extent.

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#### Ligand-induced Conformational Changes

As described herein, the unliganded receptor is in a configuration that is either inactive, has some activity or has repressor activity. Binding of agonist ligands induces conformational changes in the receptor such that the receptor becomes more active, either to stimulate or repress the expression of genes. The receptors may also have non-genomic actions. Some of the known types of changes and/or the sequelae of these are listed herein.

# Heat Shock Protein Binding

For many of the nuclear receptors ligand binding induces a dissociation of heat shock proteins such that the receptors can form dimers in most cases, after which the receptors bind to DNA and regulate transcription.

Nuclear receptors usually have heat shock protein binding domains that present a region for binding to the LBD and can be modulated by the binding of a 25 ligand to the LBD. Consequently, an extended chemical moiety (or more) from the ligand that stabilizes the binding or contact of the heat shock protein binding domain with the LBD can be designed using the computational methods described herein to produce a partial agonist or antagonist. Typically such extended chemical moieties will extend past and away from the molecular recognition domain on the ligand and 30 usually past the buried binding cavity of the ligand.

# Dimerization and Heterodimerization

With the receptors that are associated with the hsp in the absence of the ligand, dissociation of the hsp results in dimerization of the receptors. Dimerization is due to

receptor domains in both the DBD and the LBD. Although the main stimulus for dimerization is dissociation of the hsp, the ligand-induced conformational changes in the receptors may have an additional facilitative influence. With the receptors that are not associated with hsp in the absence of the ligand, particularly with the TR, ligand 5 binding can affect the pattern of dimerization/heterodimerization. The influence depends on the DNA binding site context, and may also depend on the promoter context with respect to other proteins that may interact with the receptors. A common pattern is to discourage monomer formation, with a resulting preference for heterodimer formation over dimer formation on DNA.

Nuclear receptor LBDs usually have dimerization domains that present a region for binding to another nuclear receptor and can be modulated by the binding of a ligand to the LBD. Consequently, an extended chemical moiety (or more) from the ligand that disrupts the binding or contact of the dimerization domain can be designed using the computational methods described herein to produce a partial agonist or antagonist. Typically such extended chemical moieties will extend past and away from the molecular recognition domain on the ligand and usually past the buried binding cavity of the ligand.

#### **DNA** Binding

20 In nuclear receptors that bind to hsp, the ligand-induced dissociation of hsp with consequent dimer formation allows, and therefore, promotes DNA binding. With receptors that are not associated (as in the absence of ligand), ligand binding tends to stimulate DNA binding of heterodimers and dimers, and to discourage monomer binding to DNA. However, ligand binding to TR, for example, tends to 25 decrease dimer binding on certain DNA elements and has minimal to no effect on increasing heterodimer binding. With DNA containing only a single half site, the ligand tends to stimulate the receptor's binding to DNA. The effects are modest and depend on the nature of the DNA site and probably on the presence of other proteins that may interact with the receptors. Nuclear receptors usually have DBDs that 30 present a region for binding to DNA and this binding can be modulated by the binding of a ligand to the LBD. Consequently, an extended chemical moiety (or more) from the ligand that disrupts the binding or contact of the DBD can be designed using the computational methods described herein to produce a partial agonist or antagonist. Typically such extended chemical moieties will extend past and away from the

molecular recognition domain on the ligand and usually past the buried binding cavity of the ligand.

# Repressor Binding

Receptors that are not associated with hsp in the absence of ligand frequently act as transcriptional repressors in the absence of the ligand. This appears to be due, in part, to transcriptional repressor proteins that bind to the LBD of the receptors. Agonist binding induces a dissociation of these proteins from the receptors. This relieves the inhibition of transcription and allows the transcriptional transactivation functions of the receptors to become manifest.

# Transcriptional Transactivation Functions

Ligand binding induces transcriptional activation functions in two basic ways.

The first is through dissociation of the hsp from receptors. This dissociation, with

15 consequent dimerization of the receptors and their binding to DNA or other proteins in the nuclear chromatin allows transcriptional regulatory properties of the receptors to be manifest. This may be especially true of such functions on the amino terminus of the receptors.

The second way is to alter the receptor to interact with other proteins involved 20 in transcription. These could be proteins that interact directly or indirectly with elements of the proximal promoter or proteins of the proximal promoter. Alternatively, the interactions could be through other transcription factors that themselves interact directly or indirectly with proteins of the proximal promoter. Several different proteins have been described that bind to the receptors in a ligand-25 dependent manner. In addition, it is possible that in some cases, the ligand-induced conformational changes do not affect the binding of other proteins to the receptor, but do affect their abilities to regulate transcription.

Nuclear receptors or nuclear receptor LBDs usually have activation domains modulated in part by a co-activator/co-repressor system that coordinately functions to 30 present a region for binding to DNA, and can be modulated by the binding of a ligand to the LBD. Consequently, an extended chemical moiety (or more) from the ligand that disrupts the binding or contact of the activation domain with co-activator and/or co-repressor can be designed using the computational methods described herein to produce a partial agonist or antagonist. For instance, an agonist can be designed

and/or selected which (1) blocks binding and/or dissociates co-repressor, and/or (2) promotes binding and/or association of a co-activator. An antagonist can be designed which (1) promotes binding and/or association of co-repressor, and/or (2) promotes binding and/or association of co-activator. Ratios of agonists and antagonists may be 5 used to modulate transcription of the gene of interest. Selection can be accomplised in binding assays that screen for ligands having the desired agonist or antagonist properties, including such ligands which induce conformational changes as decribed below. Suitable assays for such screening are described herein and in Shibata, H., et al. (Recent Prog. Horm. Res. 52:141-164 (1997)); Tagami, T., et al. (Mol. Cell Biol. 10 17(5):2642-2648 (1997)), Zhu, XG., et al. (J. Biol. Chem. 272(14):9048-9054 (1997)); Lin, B.C., et al. (Mol. Cell Biol. 17(10):6131-6138 (1997)); Kakizawa, T., et al. (J. Biol. Chem. 272(38):23799-23804 (1997)); and Chang, K. H., et al. (Proc. Natl. Acad. Sci. USA 94(17):9040-9045 (1997)). Typically such extended chemical moieties will extend past and away from the molecular recognition domain on the 15 ligand and usually past the buried binding cavity of the ligand and in the direction of the activation domain, which is often a helix as seen in the three dimensional model shown in the FIGURES in two dimensions on paper or more conveniently on a computer screen.

# 20 Ligand-Induced Conformational Change

Plasma proteins bind hormones without undergoing a conformational change through a static binding pocket formed between monomers or domains. For example, the tetrameric thyroid-binding plasma protein transthyretin forms a solvent-accessible hormone-binding channel at the oligomer interface. The structure of the protein is unchanged upon binding hormone with respect to the appearance of a buried binding cavity with a ligand bound.

However, the structural role for a ligand bound to a nuclear receptor LBD, like rat TR-α LBD, predicts that the receptor would differ in the bound and unbound states. In the absence of hormone, the receptor would possess a cavity at its core, 30 uncharacteristic of a globular protein. A ligand (e.g. hormone) completes the hydrophobic core of the active receptor after it binds to the nuclear receptor. Ligand binding by the receptor is a dynamic process, which regulates receptor function by inducing an altered conformation.

An exact description of the hormone-induced conformational changes requires comparison of the structures of the liganded and the unliganded TR. The structure of the unliganded human RXRα may substitute as a model for the unliganded TR. The rat TR-α LBD and human RXRα LBDs adopt a similar fold, and it is likely that the structural similarity extends to the conformational changes after ligand binding.

There are three major differences between the two structures, which indeed appear to be the result of ligand binding. First, the bound rat TR-α LBD structure is more compact, with the hormone tightly packed within the hydrophobic core of the receptor. By contrast, the unliganded human RXRα LBD contains several internal hydrophobic cavities. The presence of such cavities is unusual in folded proteins, and is likely a reflection of the unliganded state of the receptor. Two of these cavities were proposed as possible binding sites for 9-cis retinoic acid, though these multiple sites only partly overlap with the single buried binding cavity observed in the liganded rat TR-α LBD.

15 The second difference involves H11 in the rat TR-α LBD, which contributes part of the hormone binding cavity. H11, continuous in the rat TR-α LBD is broken at Cys 432 in the RXR, forming a loop between H10 and H11 in the hRXRα. This residue corresponds to His381 in the TR, which provides a hydrogen bond to the outer ring hydroxyl of the ligand. Furthermore, the hormone binding cavity occupied by ligand in the rat TR-α LBD is interrupted in the hRXRα by the same loop, forming an isolated hydrophobic pocket in the RXR with H6 and H7. In the bound rat TR-α LBD, the corresponding helices H7 and H8 are contiguous with the binding pocket, and enclose the hormone binding cavity from below.

The third difference between the two receptors is the position of the C-25 terminal activation domain. While the C-terminal activation domain forms α-helices in both receptors, the domain in the rat TR-α LBD follows a proline-rich turn, and lies against the receptor to contribute part of the binding cavity. In contrast, the activation domain in the unliganded hRXRα, is part of a longer helix which projects into the solvent.

These differences lead to a model for an alternate conformation of the TR LBD assumed in the absence of ligand. In the unliganded TR, the subdomain of the receptor surrounding the hormone binding cavity is loosely packed, with the binding cavity occluded by a partly unstructured H11 providing a partial core for the receptor.

Upon binding hormone, residues which form a coil in the unbound receptor engage the ligand, and continues H11. The ordering of H11 could unblock the hydrophobic cavity, allowing H7 and H8 to interact with hormone. The extended hydrophobic cavity then collapses around the hormone, generating the compact bound 5 structure.

It is possible to predict ligand-induced conformational changes in the C-terminal activation domain that rely, in part, on an extended structure in the unliganded TR that repacks upon ligand binding. The ligand- induced conformation change can be subtle since the amino acid sequence of the rat TR-α in the turn (393-10 PTELFPP-399) significantly reduces the propensity of the peptide chain of the rat TR-α to form an α-helix and therefore repacking can be accomplished with a minor change in volume.

After the ligand-induced conformational change occurs, it is likely that the conformation of the C-terminal activation domain in the bound structure changes 15 packing compared to the unbound form of the receptor. Binding of the ligand improves the stability of the activation domain. The activation domain packs loosely even in the bound structure, as measured by the distribution of packing interactions for the entire LBD. The packing density for the activation domain, defined as the number of atoms within 4.5Å, is 1.5 standard deviations below the mean. For 20 comparison, another surface helix, H1, is 0.5 standard deviations below the mean and the most poorly packed part of the structure, the irregular coil from residues Ile196-Asp206, is 2.0 standard deviations below the mean. Moreover, the majority of packing contacts for the C-terminal domain in the bound receptor are provided either by residues which interact with ligand, such as His381, or by the ligand itself. The 25 conformation of these residues can be expected to be different in the bound and unbound receptors, and by extension the conformation of C-terminal activation domain which relies upon these interactions. Without the stabilization provided by a bound ligand, it is likely that the C-terminal activation domain is disordered prior to hormone binding.

The interrelation of ligand-induced conformational changes is evident as described herein. For example, His381 from H11 and Phe405 from H12 interact in the bound structure to provide a specific hydrogen bond to the phenolic hydroxyl. The

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WO 99/26966 PCT/US98/25296

ligand-induced changes which affect H11 and H12 are reinforcing, and lead to the formation of the compact, bound state.

Comparison of the TR- $\alpha$  and TR- $\beta$  LBD structures shows similar packing of the helices when complexed with the ligand Triac.

# COMPUTATIONAL METHODS USING THREE DIMENSIONAL MODELS AND EXTENSIONS OF LIGANDS

The three-dimensional structure of the liganded TR receptor is unprecedented, and will greatly aid in the development of new nuclear receptor synthetic ligands, 10 such as thyroid receptor antagonists and improved agonists, especially those that bind selectively to one of the two TR isoforms ( $\alpha$  or  $\beta$ ). In addition, this receptor superfamily is overall well suited to modern methods including three-dimensional structure elucidation and combinatorial chemistry such as those disclosed in EP 335 628, U.S. patent 5,463,564, which are incorporated herein by reference. Structure 15 determination using X-ray crystallography is possible because of the solubility properties of the receptors. Computer programs that use crystallography data when practicing the present invention will enable the rational design of ligand to these receptors. Programs such as RASMOL can be used with the atomic coordinates from crystals generated by practicing the invention or used to practice the invention by 20 generating three dimensional models and/or determining the structures involved in ligand binding. Computer programs such as INSIGHT and GRASP allow for further manipulation and the ability to introduce new structures. In addition, high throughput binding and bioactivity assays can be devised using purified recombinant protein and modern reporter gene transcription assays described herein and known in the art in 25 order to refine the activity of a CDL.

Generally the computational method of designing a nuclear receptor synthetic ligand comprises two steps:

- determining which amino acid or amino acids of a nuclear receptor LBD interacts with a first chemical moiety (at least one) of the ligand using a three
   dimensional model of a crystallized protein comprising a nuclear receptor LBD with a bound ligand, and
  - 2) selecting a chemical modification (at least one) of the first chemical moiety to produce a second chemical moiety with a structure to either decrease or increase an

interaction between the interacting amino acid and the second chemical moiety compared to the interaction between the interacting amino acid and the first chemical moiety.

As shown herein, interacting amino acids form contacts with the ligand and 5 the center of the atoms of the interacting amino acids are usually 2 to 4 angstroms away from the center of the atoms of the ligand. Generally these distances are determined by computer as discussed herein and in McRee 1993, however distances can be determined manually once the three dimensional model is made. Examples of interacting amino acids are described in Appendix 2. See also Wagner et al., Nature 10 378(6558):670-697 (1995) for stereochemical figures of three dimensional models. More commonly, the atoms of the ligand and the atoms of interacting amino acids are 3 to 4 angstroms apart. The invention can be practiced by repeating steps 1 and 2 to refine the fit of the ligand to the LBD and to determine a better ligand, such as an agonist. As shown in the FIGURES the three dimensional model of TR can be 15 represented in two dimensions to determine which amino acids contact the ligand and to select a position on the ligand for chemical modification and changing the interaction with a particular amino acid compared to that before chemical modification. Structural comparison of LBD isoforms complexed with the same or similar ligand permit identification of fiducial and adjustable amino acids that can be 20 exploited in designing isoform-specific ligands through chemical modification. "Fiducial" refers to amino acids that form rigid features of the ligand binding cavity. "Adjustable" refers to amino acids that form less rigid features of the ligand binding cavity. The chemical modification may be made using a computer, manually using a two dimensional representation of the three dimensional model or by chemically 25 synthesizing the ligand. The three dimensional model may be made using Appendix 2 and the FIGURES. As an additional step, the three dimensional model may be made using atomic coordinates of nuclear receptor LBDs from crystallized protein as known in the art, see McRee 1993 referenced herein.

The ligand can also interact with distant amino acids after chemical modification of the ligand to create a new ligand. Distant amino acids are generally not in contact with the ligand before chemical modification. A chemical modification can change the structure of the ligand to make as new ligand that interacts with a distant amino acid usually at least 4.5 angstroms away from the ligand. Often distant

amino acids will not line the surface of the binding cavity for the ligand, as they are too far away from the ligand to be part of a pocket or surface of the binding cavity.

The interaction between an atom of a LBD amino acid and an atom of an LBD ligand can be made by any force or attraction described in nature. Usually the 5 interaction between the atom of the amino acid and the ligand will be the result of a hydrogen bonding interaction, charge interaction, hydrophobic interaction, van der waals interaction or dipole interaction. In the case of the hydrophobic interaction it is recognized that this is not a per se interaction between the amino acid and ligand, but rather the usual result, in part, of the repulsion of water or other hydrophilic group 10 from a hydrophobic surface. Reduction or enhancment of the interaction of the LBD and a ligand can be measured by standard binding procedures, calculating or testing binding energies, computationally or using thermodynamic or kinetic methods as known in the art.

Chemical modifications will often enhance or reduce interactions of an atom of a LBD amino acid and an atom of an LBD ligand. Steric hinderance will be a common means of changing the interaction of the LBD binding cavity with the activation domain. Chemical modifications are preferably introduced at C-H, C- and C-OH position in ligands, where the carbon is part of the ligand structure which remains the same after modification is complete. In the case of C-H, C could have 1, 2 or 3 hydrogens, but usually only one hydrogen will be replaced. The H or OH are removed after modification is complete and replaced with the desired chemical moiety.

Because the thyroid receptor is a member of the larger superfamily of hormone-binding nuclear receptors, the rules for agonist and antagonist development will be recognized by one skilled in the art as useful in designing ligands to the entire superfamily. Examining the structures of known agonists and antagonists of the estrogen and androgen receptors supports the generality of antagonist mechanism of action as shown in FIG. 10.

The overall folding of the receptor based on a comparison of the reported structure of the unliganded RXR and with amino acid sequences of other superfamily members reveals that the overall folding of receptors of the superfamily is similar. Thus, it is predicted from the structure that there is a general pattern of folding of the nuclear receptor around the agonist or antagonist ligand.

The three dimensional structure of a nuclear receptor with a ligand bound leads to the nonobvious observation that a nuclear receptor folds around agonist ligands, as the binding cavity fits the agonist, especially the agonist's molecular recognition domain, and antagonists commonly have chemical structures that extend 5 beyond the ligand, especially the agonist, and would prohibit folding of the receptor around the ligand to form a buried binding cavity or other groups that have the same effect. The location of the extension could affect the folding in various ways as indicated by the structure. Such extensions on antagonists are shown in FIG. 10 for various receptors and compared to the corresponding agonist.

10 For example, an extension towards the carboxy-terminal activation helix affects the packing/folding of this helix into the body of the receptor. This in turn can affect the ability of this portion of the nuclear receptor to interact with other proteins or other portions of the receptor, including transcriptional transactivation functions on the opposite end of the linear receptor, or the receptor's amino terminus that may 15 interact directly or indirectly with the carboxy-terminal transactivation domain (including helix 12). Extensions in this direction can also affect the packing of helix 11 of TR (or its analogous helix in nuclear receptors) into the body of the receptor and selectively affect dimerization and heterodimerization of receptors. An extension pointing towards helix 1 can affect the relationship of the DNA binding domain and 20 hinge regions of the receptors with the ligand binding domain and selectively or in addition affect the receptors' binding to DNA and/or interactions of receptors with proteins that interact with this region of the receptor. Other extensions towards helix 11 can be made to affect the packing of this helix and helices 1 and 10 and thereby homo- and hetero-dimerization. Such chemical modifications can be assessed using 25 the computational methods described herein. It is also possible that, in some cases, extensions may protrude through the receptor that is otherwise completely or incompletely folded around the ligand. Such protruding extensions could present a steric blockade to interactions with co-activators or other proteins.

The three dimensional structure with the ligand buried in the binding cavity immediately offers a simple description of a nuclear receptor that has a binding cavity that contains hinges and a lid, composed of one or more structural elements, that move to accommodate and surround the ligand. The ligand to TR can be modified on specific sites with specific classes of chemical groups that will serve to leave the lid and hinge region in open, partially open or closed states to achieve partial agonist or

antagonist functions. In these states, the biological response of the TR is different and so the structure can be used to design particular compounds with desired effects.

Knowledge of the three-dimensional structure of the TR-T<sub>3</sub> complex leads to a general model for agonist and antagonist design. An important novel feature of the structural data is the fact that the T<sub>3</sub> ligand is completely buried within the central hydrophobic core of the protein. Other ligand-receptor complexes belonging to the nuclear receptor superfamily will have a similarly buried ligand binding site and therefore this model will be useful for agonist/antagonist design for the entire superfamily.

When design of an antagonist is desired, one needs either to preserve the important binding contacts of natural hormone agonist while incorporating an "extension group" that interferes with the normal operation of the ligand-receptor complex or to generate the requisite binding affinity through the interactions of the extensions with receptor domains.

15 The model applied to antagonist design and described herein is called the "Extension Model." Antagonist compounds for nuclear receptors should contain the same or similar groups that facilitate high-affinity binding to the receptor, and in addition, such compounds should contain a side chain which may be large and/or polar. This side chain could be an actual extension, giving it bulk, or it could be a 20 side group with a charge function that differs from the agonist ligand. For example, substitution of a CH<sub>3</sub> for CH<sub>2</sub>OH at the 21-position, and alteration at the 11-position from an OH group to a keto group of cortisol generates glucocorticoid antagonist activity (Robsseau, G.G., et. al., J. Mol. Biol. 67:99-115 (1972)). However, in most cases effective antagonists have more bulky extensions. Thus, the antiglucocorticoid 25 (and antiprogestin) RU486 contains a bulky side group at the 11-position (Horwitz, K.B. Endocrine Rev. 13:146-163 (1992)). The antagonist compound will then bind within the buried ligand binding site of the receptor with reasonably high affinity (100 nM), but the extension function will prevent the receptor-ligand complex from adopting the necessary conformation needed for transcription factor function. The 30 antagonism (which could be in an agonist or antagonist) may manifest itself at the molecular level in a number of ways, including by preventing receptor homo/heterodimer formation at the HRE, by preventing coactivator binding to receptor monomers, homodimers or homo/heterodimers, or by a combination of these effects which otherwise prevent transcription of hormone responsive genes mediated

by ligand-induced effects on the HRE. There are several antagonist compounds for nuclear receptors in the prior art (see also Horwitz, K.B., Endocrine Rev. 13:146-163 (1992), Raunnaud J.P. et. al., J. Steroid Biochem. 25:811-833 (1986), Keiel S., et. al., Mol. Cell. Biol. 14:287-298 (1994) whose antagonist function can be explained by the extension hypothesis. These compounds are shown in FIG. 10 along with their agonist counterparts. Each of these antagonists contains a large extension group attached to an agonist or agonist analogue core structure. Importantly, these antagonist compounds were discovered by chance and not designed with a structure-function hypothesis such as the extension principle.

One method of design of a thyroid antagonist using the extension hypothesis is provided below as a teaching example. The three-dimensional structure of the TR-α Dimit complex combined with structure-activity data published in the prior art, especially those reference herein, can be used to establish the following ligand-receptor interactions which are most critical for high-affinity ligand binding.

15 A physical picture of these interactions is shown in FIG. 6. The figure describes the isolated essential contacts for ligand binding. Because the ligand is buried in the center of the receptor, the structural spacing between these isolated interactions is also important. Thus, our present knowledge of this system dictates that, for this example, a newly designed ligand for the receptor must contain a thyronine structural skeleton, or two substituted aryl groups joined by a one-atom spacer.

The general structure for an antagonist designed by the extension hypothesis is exemplified in the following general description of the substituents of a TR antagonist (referring to Formula 1): R<sub>1</sub> can have anionic groups such as a carboxylate, phosphonate, phosphate, sulfate or sulfite and is connected to the ring with a 0 to 3 atom linker, comprising one or more C, O, N, S atoms, and preferably a 2 carbon linker. Such R<sub>1</sub> can be optionally substituted with an amine (e.g. -NH<sub>2</sub>). R<sub>3</sub> and R<sub>5</sub> are small hydrophobic groups such as -Br, -I, or -CH<sub>3</sub>. R<sub>3</sub> and R<sub>5</sub> can be the same substituents or different. R<sub>3</sub>' can be a hydrophobic group that may be larger than those of R<sub>3</sub> and R<sub>5</sub>, such as -I, -CH<sub>3</sub>, -isopropyl, -phenyl, -benzyl, 5 and 6 ring heterocycles. R<sub>4</sub>' is a group that can participate in a hydrogen bond as either a donor or acceptor. Such groups include -OH, -NH<sub>2</sub>, and -SH. R<sub>5</sub>' is an important extension group that makes this compound an antagonist. R<sub>5</sub>' can be a long chain alkyl (e.g. 1 to 9 carbons, straight chain or branched), aryl (benzyl, phenyl and substituted benzyl

and phenyl rings (e.g. with halogen, alkyl (1 and 5 carbons) and optionally connected to the ring by a -CH<sub>2</sub>-), heterocycle (e.g. 5 or 6 atoms, preferably 5 carbons and 1 nitrogen, or five carbons), which can optionally include polar (e.g. -OH, -NH<sub>2</sub>, and -SH), cationic (e.g. -NH<sub>3</sub>, N(CH)<sub>3</sub>), or anionic (carboxylate, phosphonate, phosphate or sulfate) groups. R<sub>5</sub>' can also be a polar (e.g. -OH, -NH<sub>2</sub>, and -SH), cationic (e.g. -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>), and anionic (carboxylate, phosphonate, phosphate or sulfate) groups. X is the spacer group that appropriately positions the two aromatic rings. This group is usually a one-atom spacer, such as O, S, SO, SO<sub>2</sub>, NH, NZ where Z is an alkyl, CH<sub>2</sub>, CHOH, CO, C(CH<sub>3</sub>)OH, and C(CH<sub>3</sub>)(CH<sub>3</sub>). X also may be NR<sub>7</sub>, CHR<sub>7</sub>, CR<sub>7</sub>, R<sub>7</sub>, where R<sub>7</sub>, is an alkyl, aryl or 5- or 6-membered heterocyclic aromatic. R<sub>2</sub>, R<sub>6</sub>, R<sub>2</sub>' and R<sub>6</sub>' can be -F, and -Cl and are preferably H.

A TR ligand can also be described as a substituted phenylated 3,5 diiodo tyrosine with substituted R<sub>5</sub>' and R<sub>3</sub>' groups. R<sub>5</sub>' can be a long chain alkyl (e.g. 4 to 9 carbons, straight chain or branched), aryl (benzyl, phenyl and substituted benzyl and 15 phenyl rings (e.g with halogen, alkyl (1 and 5 carbons) and optionally connected to the ring by a -CH<sub>2</sub>-), heterocycle (e.g. 5 or 6 atoms, preferably 5 carbons and 1 nitrogen, or five carbons), which can optionally include polar (e.g. -OH, -NH<sub>2</sub>, and -SH), cationic (e.g. -NH<sub>3</sub>, N(CH)<sub>3</sub>), or anionic (carboxylate, phosphonate, phosphonate or sulfate) groups. R<sub>5</sub>' can also be a polar (e.g. -OH, -NH<sub>2</sub>, and -SH), cationic (e.g. - NH<sub>3</sub>, N(CH)<sub>3</sub>), and anionic (carboxylate, phosphonate, phosphate or sulfate) groups. R<sub>3</sub>' can be -IsoPr, halogen, -CH<sub>3</sub>, alkyl (1 to 6 carbons) or aryl (benzyl, phenyl and substituted benzyl and phenyl rings (e.g with halogen, alkyl (1 and 5 carbons) and optionally connected to the ring by a -CH<sub>2</sub>-), heterocycle (e.g. 5 or 6 atoms, preferably 5 carbons and 1 nitrogen, or five carbons), which can optionally include polar (e.g. - 25 OH, -NH<sub>2</sub>, and -SH), cationic (e.g. -NH<sub>3</sub>, N(CH)<sub>3</sub>), or anionic (carboxylate, phosphonate, phosphate or sulfate) groups.

A TR antagonist can also be a modified T<sub>3</sub> agonist (having a biphenyl structure) wherein R<sub>5</sub>' is alkyl, aryl, 5- or 6-membered heterocyclic aromatic, heteroalkyl, heteroaryl, arylalkyl, heteroaryl alkyl, polyaromatic, polyheteroaromatic, 30 polar or charged groups, wherein said R<sub>5</sub>' may be substituted with polar or charged groups. The R<sub>5</sub>' groups are defined, as described herein.

Using these methods the ligands of this example preferably have the following properties:

1. The compounds should bind to the TR with high affinity (for example 100 nM).

- 2. The compounds should bind the receptor in the same basic orientation as the natural hormone.
- 5 3. The extension group  $R_5$ ' should project toward the activation helix (C-terminal helix) of the receptor.
  - 4. The appropriate substituent at  $R_5$ ' should perturb the activation helix from its optimal local structure needed for mediating transcription.

Antagonists may also be designed with multiple extensions in order to block 10 more than one aspect of the folding at any time.

TR ligands (e.g. super agonists) can be designed (and synthesized) to enhance the interaction of at least one amino acid with at least one chemical moiety on the ligand's molecular recognition domain. One method is to enhance the charge and polar interactions by replacing the carboxylate of T<sub>3</sub> (R<sub>1</sub> position) with phosphonate, phosphate, sulfate or sulfite. This enhances the interaction with Arg 262, Arg 266 and Arg 228. The interaction of at least one amino acid with at least one chemical moiety on the ligand's molecular recognition domain can also be enhanced by increasing the size of R<sub>1</sub> group to fill the space occupied by water when Dimit is bound (referring to R<sub>1</sub>). Preferably the group has a complementary charge and hydrophobicity to the binding cavity.

Another way of improving the interaction of at least one amino acid with at least one chemical moiety on the ligand's molecular recognition domain is to restrict the conformation of the dihedral angle between the two phenyl rings of the thyronine ligand in solution. In solution the planes of two phenyl rings are orthogonal where the dihedral angle is 90°. In the TR Dimit structure, the dihedral angle is close to 60°. A TR ligand design that fixes the angle between the two phenyl rings will lead to tighter binding. Such a ligand may be made by connecting the R<sub>6</sub>' and the R<sub>5</sub> positions of a thyronine or a substituted thyronine-like biphenyl. The size of the cyclic connection can fix the angle between the two phenyl rings. Referring specifically to Formula 1, 30 the following cyclic modifications are preferred: 1) R<sub>5</sub> is connected to R<sub>6</sub>', 2) R<sub>3</sub> is connected to R<sub>2</sub>' or 3) R<sub>5</sub> is connected to R<sub>6</sub>' and R<sub>3</sub> is connected to R<sub>2</sub>'. The connections can be made by an alkyl or heteroalkyl chain having between 1 to 6 atoms and preferably from 2 to 4 carbon atoms or other atoms. Any position of the heteroalkyl chain can be N, O, P or S. The S and P heteroatoms along said heteroalkyl

chain are in any of their possible oxidative states. The N heteroatom or any carbon along the alkyl or heteroalkyl chain may have one or more Z substituents, wherein Z is alkyl, heteroalkyl, aryl, heteroaryl, 5- or 6-membered heterocyclic aromatic. These compounds can be claimed with the proviso that Formula 1 does not include any prior art compound as of the priority filing date of this application.

The interaction of at least one amino acid with at least one chemical moiety on the ligand's molecular recognition domain can also be enhanced by selecting a chemical modification that fills the unfilled space between a TR ligand and the LBD in the area of the bridging oxygen (such as in T3, Triac or Dimit). Thus, a slighter larger moiety that replaces the ether oxygen can enhance binding. Such a linker may be a mono- or geminal- disubstituted carbon group. A group approximately the same size as oxygen but with greater hydrophobicity is preferred as well as small, hydrophobic groups for the disubstituted carbon.

Compounds of Formula I or derivatives thereof that modulate TR activity also may be designed and selected to interact with a conformationally constrained structural feature of a TR LBD that is conserved among TR LBD isoforms to increase TR-specific selectivity. Conserved structural features of a TR LBD include residues found in equivalent positions of TR LBD isoforms which interact with a conserved structural feature of a compound comprising the biphenyl scaffold (φ-X-φ) or a single phenyl scaffold (φ-X or X-φ) of Formula I. Conformationally constrained structural features of a TR LBD include residues that have their natural flexible conformations fixed by various geometric and physical-chemical constraints, such as local backbone, local side chain, and topological constraints. These types of constraints are exploited to restrict positioning of atoms involved in receptor-ligand recognition and binding.

25 For example, comparison of atomic models of TR LBD isoforms bound to thyronine and thyronine-like ligands reveal that certain residues which contact the ligands are restricted to particular topological shapes and angles of rotation about bonds. These

30 Ile275 and Phe455, respectively.
Selectivity imparted by conformationally constrained features of both the receptor and compound are of particular interest. For example, compounds of

Formula I comprising constrained cyclic carbons and substituent groups that interact

include Met259, Leu276, Leu292, His381, Gly290, Ile221, and Phe401 of TR-α. The corresponding positions in TR-β include Met313, Leu330, Leu346, His435, Gly344,

with a constrained feature of a TR LBD can be exploited to further increase binding specificity while reducing the potential for cross-over interaction with other receptors. These include hydrophobic and/or hydrophilic contacts between constrained residues of a TR LBD and atomic groups of the following constituents of the compound in 5 reference to Formula I: (i) the biphenyl rings; (ii) the R<sub>3</sub>-substituent; (iii) the R<sub>3</sub>'-substituent; and (iv) the R<sub>4</sub>'-substituent.

For example, contacts to the phenyl moiety comprising the R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>5</sub> and R<sub>6</sub> substituents, i.e., the ring proximal to the polar pocket (the "inner ring"), include a cycle carbon atom that interacts with an atom of a hydrophobic residue of a TR LBD, such as a carbon and oxygen atom of Met259 and a carbon atom of Leu276 of TR-α, or Met313 and Leu330 of TR-β, where the cycle carbon is about 3.0 to 4.0A from the atom of the hydrophobic group. For example, comparison of TR-α complexed with T3 and TR-β complexed with GC-1 reveals the following conserved inner ring contacts:

15				
	Ligand	TR LB	D	
	T3/Atom	TR-α Residue	Atom	Distance
	C11	Met259	С	3.95
	C11	Met259	0	3.59
20	C11	Met259	CB	3.77
	C7	Leu276	CD2	3.80
	C9	Leu276	CD2	3.70
		•		
	GC1/Atom	TR-β Residue	Atom	Distance
25	C11	Met313	С	3.85
	C11	Met313	O	3.41
	C11	Met313	CB	3.79
	C7	Leu330	CD2	3.56
	C9	Leu330	CD2	3.63
30				

Contacts to the phenyl moiety comprising the R<sub>2</sub>', R<sub>3</sub>', R<sub>4</sub>', R<sub>5</sub>' and R<sub>6</sub>' substituents, i.e., the ring distal to the polar pocket (the "outer ring"), include a cyclic carbon atom that interacts with an atom of a hydrophobic residue of a TR LBD, such 35 as a carbon atom of Leu292 of TR-α, or Leu346 of TR-β, where the cyclic carbon atom is about 3.0 to 4.0A from the atom of the hydrophobic residue. For example, comparison of TR-α complexed with T3 and TR-β complexed with GC-1 reveals the following conserved outer ring contacts:

	<u>Ligand</u>	TR LB	TR LBD		
	T3/Atom	TR-α Residue	Atom	Distance	
	C6	Leu292	CD2	3.58	
	C8	Leu292	CD2	3.50	
5	GC1/Atom	TR-β Residue	Atom	Distance	
	C6	Leu346	CD2	3.77	
	C8	Leu346	CD2	3.80	

Contacts to the R<sub>3</sub>-substituent include an atom that interacts with a carbon atom of a hydrophobic residue of a TR LBD, such as Ile221 of TR-α, or Ile275 of TR-β, where the R<sub>3</sub>-substituent atom is about 3.0 to 4.0A from the carbon atom of the hydrophobic residue. For example, comparison of TR-α complexed with T3 and TR-β complexed with GC-1 reveals the following conserved R<sub>3</sub>-substituent contacts:

15

	<u>Ligand</u>	TR LB	TR LBD	
	T3/Atom I1	TR-α Residue Ile221	Atom CG1	Distance 4.01
20	GC1/Atom C19	TR-β Residue Ile275	Atom CG1	Distance 3.98

Contacts to the R<sub>3</sub>'-substituent include an atom that interacts with an atom of a hydrophobic or hydrophilic residue of a TR LBD, such as an oxygen atom of Gly290 25 of TR-α, or Gly344 of TR-β, where the R<sub>3</sub>'-substituent atom is about 3.0 to 4.0A from the atom of the hydrophobic or hydrophilic residue. For example, comparison of TR-α complexed with T3 and TR-β complexed with GC-1 reveals the following conserved R<sub>4</sub>'-substituent, phenolic hydroxyl contacts:

30	<u>Ligand</u>	TR LBD	
	T3/Atom	TR-α Residue Ator	n Distance
	I2	Gly290 O	3.50
2.5	GC1/Atom	TR-β Residue Aton	n Distance
	C18	Gly344 O	3.60

35

Contacts to the  $R_4$ '-substituent comprising a phenolic hydroxyl include carbon and oxygen atoms that interact with a hydrophobic or hydrophilic residue of a TR LBD, such as a carbon and nitrogen atom of His381 of TR- $\alpha$ , or His435 of TR- $\beta$ ,

where the  $R_4$ '-substituent atom is about 2.0 to 4.0A from an atom of the hydrophobic or hydrophilic residue. For example, comparison of TR- $\alpha$  complexed with T3 and TR- $\beta$  complexed with GC-1 reveals the following conserved  $R_4$ '-substituent, phenolic hydroxyl contacts:

5			
	Ligand	TR LBD	_
	T3/Atom	TR-a Residue Atom	Distance
	C10	His381 CD2 3.97	
	O1	His381 CD2 3.39	
10	O1	His381 CE1 3.82	
	C8	His381 NE2 3.47	
	C10	His381 NE2 3.55	
	O1	His381 NE2 2.70	
	GC1/Atom	TR-β Residue Atom	Distance
15	C10	His435 CD2 3.89	
	O1	His435 CD2 3.64	
	01	His435 CE1 3.79	
	C8	His435 NE2 3.44	
	C10	His435 NE2 3.33	
20	O1	His435 NE2 2.77	

Contacts to the R<sub>4</sub>'-substituent also may include an atom that interacts with a carbon atom of a hydrophobic residue of a TR LBD, such as Phe401 of TR-α, or 25 Phe455 of TR-β, for defining agonist activity, i.e., proper presentation of helix-12 (H12) of the TR LBD following ligand binding. The R<sub>4</sub>'-substituent atom is about 3.0 to 4.0Å from the carbon atom of the hydrophobic group. For example, comparison of TR-α complexed with T3 and TR-β complexed with GC-1 reveals the following conserved R<sub>4</sub>'-substituent contacts:

30				
	Ligand	TR LB	D	-
	T3/Atom	TR-α Residue	Atom	Distance
	O1 .	Phe401	CE1	3.52
	O1	Phe401	CZ	3.32
35	GC1/Atom	TR-β Residue	Atom	Distance
	O1	Phe455	CE1	3.40
	O1	Phe455	CZ	3.22

40 Comparison of atomic models of TR LBD isoforms complexed with the same and/or different ligands therefore facilitates the identification of new compounds that

fit spacially and preferentially into a TR LBD. Modeling, comparison of TR-ligand overlays, and comparison of TR LBD isoforms also permit identification of conformationally conserved structural features of TR LBD/ligand contacts. Exploiting conformational constraints of the LBD-ligand interaction identified by such methods therefore improves the design and identification of new compounds having increased selectivity for binding a particular type of nuclear receptor, such as TR.

# TR-α AND TR-β SELECTIVITY FOR THE THYROID HORMONE RECEPTOR

Using the method described herein ligands can be designed that selectively bind to the alpha more than the beta TR or vice versa. The X-ray crystallographic structure of the rat TR-α LBD provides insight into design of such ligands.

The three dimensional structure reveals that the major difference between the TR-α and TR-β in the ligand binding cavity resides in amino acid Ser 277 (with the 15 side group -CH<sub>2</sub>OH) in the rat TR-α and whose corresponding residue is 331, asparagine (with the side group -CH<sub>2</sub>CONH<sub>2</sub>), in the human TR-β. The side chain in human TR-β is larger, charged and has a different hydrogen bonding potential, which would allow the synthesis of compounds that discriminate between this difference. The Ser277 (Asn331 in TR-β) forms part of the polar pocket of the TR LBD, indicating that for TR-α versus TR-β discrimination, ligands can be designed to contain chemical modification of the R1-substitutent with reference to Formula I that exploit this difference.

For example, in the complex of TR-α with Triac, Ser277 does not participate in ligand binding. The absence of a role for Ser277 (Asn331 in beta) is consistent 25 with the equal affinity of Triac for the alpha and beta isoforms, and indirectly supports the contention that alpha/beta selectivity resides in the amino acid substitution Ser277 to Asn331 and its interaction with Arg228. The effect of the amino acid substitution is further evident when the interactions of Asn331 and Arg282 in the structures of the TR-β LBD complexed with GC-1 or Triac are compared with those of Ser277 and Arg228 in the TR-α LBD. In the complex with GC-1, Asn331 forms a hydrogen bond to Arg282, which in turn forms a hydrogen bond with the carboxylate of GC-1, a pattern that resembles the interactions of Ser277 and Arg228 in the complexes of the TR-α LBD complexed with T<sub>3</sub> or Triac.

However, in the complex of TR-β with Triac, Arg282 rotates away from Asn331 and the ligand, instead forming hydrogen bonds to residues Thr287 and Asp291 of H3. Therefore, differences exist between the two isoforms in the conformation of the polar pocket, depending on the nature of the ligand R<sub>1</sub>-substitutent, indicating that certain substituents may interact preferentially with the conformation of a given isoform.

Comparing overlays of various ligands bound to the TR- $\alpha$  versus TR- $\beta$  LBDs shows the positioning of the ligand to be very similar. Surprisingly, comparison of the volume and area for the  $TR-\alpha$  and  $TR-\beta$  LBDs bound by the same or different ligands unexpectedly shows that the cubic space or volume available for 10 accommodating ligand binding by the TR- $\beta$  LBD (645 ± 28.28 Å<sup>3</sup>) is larger and more flexable than that of the TR- $\alpha$  LBD (596.25 ± 7.97 Å<sup>3</sup>) (Table 1). The volume of the ligand binding cavity for TR-α varies over a narrow range of about 8+, with a maximum difference of about 16+. In contrast, the volume of the ligand binding cavity for TR-B differs by nearly 40+ between the complexes with GC-1 and Triac. 15 There also is a difference in the volume of the ligand binding cavity when comparing the same ligand bound to TR- $\alpha$  and TR- $\beta$ . For example, TR- $\alpha$  and TR- $\beta$  complexed with Triac differ in LBD volume by about 36 Å<sup>3</sup>. Comparison of TR-α and TR-β bound to Dimit and GC-1, respectively, which ligands have similar volume/area and superpositioned architecture, show that the difference in LBD volume is about 75 Å<sup>3</sup>. 20 These differences are attributed primarily to variable movement and interaction of side chain groups with ligand substitutents of the phenyl moiety ( $\phi$ ) of the biphenyl scaffold  $(\phi - X - \phi)$  located proximal to the polar pocket, e.g.,  $R_1$ -substituents in reference to Formula I. In contrast, the volume available in the hydrophobic pocket for both the TR- $\alpha$  and TR- $\beta$  LBDs is substantially the same. For example, binding of 25 Triac to the TR-β LBD displaces the side chain of Arg 282 providing approximately 60 Å<sup>3</sup> in the polar pocket cavity, exposing the polar pocket to bulk solvent exchange. For GC-1 bound to the TR-β LBD, approximately 14 Å<sup>3</sup> is due to side chain motion of Met310, and approximately 44 Å<sup>3</sup> is due to side chain motion of Arg320, the combination of which increases the size of the polar pocket in the TR-β LBD. This 30 extra pliability also may explain the absence of ordered water in the polar pocket of TR-β LBD bound to Triac or GC-1, which is in contrast to the ordered water found in the polar pocket of TR-α LBD bound to Dimit, IpBr2 or T3.

5

WO 99/26966 PCT/US98/25296

Table 1\*

rTR-α						
	Dimit	Triac	IpBr2	T3		
TR LBD (vol $^3$ /area $^2$ )	590/456	589/440	601/474	605/472		
Ligand (volÅ <sup>3</sup> /areaÅ <sup>2</sup> )	303/314	333/326	326/330	355/346		
Complementarity	0.65	0.68	0.66	0.71		

hTR-β					
	GC-1	Triac			
TR LBD (volų/areaŲ)	665/575	625/474			
Ligand (volÅ <sup>3</sup> /areaÅ <sup>2</sup> )	294/310	333/326			
Complementarity	0.61	0.67			

\*TR LBD volume and area are reported in Angstroms measured by GRASP. Complementarity is determined as defined in Lawrence *et al.*, *J. Mol. Biol. 234*:946-950 (1993).

Residue Ser277 in TR-α and the corresponding residue Asn331 of TR-β also contribute to the volumetric differences observed in the polar pockets of these two TR isoforms. And substitution of the Asn331 of hTR-β with serine has the affect of modifying ligand binding affinity of TR-β so that it resembles that of TR-α (See Example 5). Taken together, differences in hydrogen bonding of atoms of the side chain group of Ser277 in TR-α and Asp331 in TR-β extending from the equivalent backbone position in these TR LBDs and the more restricted polar pocket of the TR-α LBD further supports the concept of designing TR LBD isoform-specific ligands 15 having substitutents that fit spacially and preferentially into the polar pocket of either the TR-α or TR-β LBDs. Exploitation of this difference provides an additional means for computational design of isoform-specific TR agonists and antagonists.

In terms of ligand design, these differences mean that for  $\beta$ -selective ligands, some or all of the following differences should be exploited:

- 20 1. The presence of a larger side chain asparagine.
  - 2. The ability of the carbonyl group on the side chain to provide a strong hydrogen bond acceptor.
  - 3. The ability of the amido group on the side chain to provide a two hydrogen bond donors.
- 25 4. Adjustment of polarity to reorganize the trapped water in the T3 pocket.
  - 5. Greater size and flexibility of the polar pocket.

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WO 99/26966 PCT/US98/25296

In terms of pharmaceutical design, these differences mean that for  $\alpha$ -selective ligands, some or all of the following differences should be exploited:

- 1. The presence of a smaller side group.
- 2. The ability of the hydroxyl on the -CH<sub>2</sub>OH side group carbonyl group on the side chain to provide a weak hydrogen donor.
  - 3. Adjustment of polarity to reorganize the trapped water in the T3 pocket.
  - 4. Smaller size and limited flexibility of the polar pocket.

In both cases these differences can be exploited in a number of ways. For example, they can also be used with a software set for construction of novel organic molecules such as LUDI from Biosym-MSI. An example of designing TR-β selective ligands is increasing the polarity of a ligand substituent located in the polar pocket of a TR LBD through addition of one or more ligand groups having a formal negative charge and/or negative dipole charge that interacts with a formal positive charge and/or positive dipole charge of a group in the polar pocket of the LBD. This exploits preferential interactions, such as with the additional positive charge contributed by Asn 331 in TR-β. Another example of a TR-β selective ligand is one that comprises one or more groups which fit spacially into the TR-β LBD polar pocket. This exploits spacial differences between TR LBD isoforms, such as the larger and more flexible 20 polar pocket of TR-β.

#### METHODS OF TREATMENT

The compounds of Formula 1 can be useful in medical treatments and exhibit biological activity which can be demonstrated in the following tests:

25 (i) the induction of mitochondrial α-glycerophosphate dehydrogenase (GPDH:EC 1.1.99.5). This assay is particularly useful since in certain species e.g. rats it is induced specifically by thyroid hormones and thyromimetics in a close-related manner in responsive tissues e.g. liver, kidney and the heart (Westerfield, W.W., Richert, D.A. and Ruegamer, W.R., *Endocrinology* (1965) 77:802). The assay 30 allows direct measurement in rates of a thyroid hormone-like effect of compounds and in particular allows measurement of the direct thyroid hormone-like effect on the heart. Other measurements included parameters such as heart rate and cardiac

enzymes including Ca<sup>++</sup> ATPase, Na<sup>++</sup>/K<sup>+</sup> ATPase, myosin isoforms and specific liver enzymes;

- (ii) the elevation of basal metabolic rate as measured by the increase in whole body oxygen consumption (see e.g., Barker et al., Ann. N. Y. Acad. Sci., (1960)
   5 86:545-562);
  - (iii) the stimulation of the rate of beating of atria isolated from animals previously dosed with thyromimetrics (see e.g., Stephan *et al.*, *Biochem. Pharmacol.* (1992) 13:1969-1974; Yokoyama *et al.*, *J. Med. Chem.*, (1995) 38:695-707);
- (iv) the change in total plasma cholesterol levels as determined using a 10 cholesterol oxidase kit (for example, the Merck CHOD iodine colorimetric kit. see also, Stephan *et al.* (1992));
- (v) the measurement of LDL (low density lipoprotein) and HDL (high density lipoprotein) cholesterol in lipoprotein fractions separated by ultracentrifugation; and p (vi) the change in total plasma triglyceride levels as 15 determined using enzymatic color tests, for example the Merck System GPO-PAP method.

The compounds of Formula 1 can be found to exhibit selective thyromimetic activity in these tests,

- (a) by increasing the metabolic rate of test animals, and raising hepatic 20 GPDH levels at doses which do not significantly modify cardiac GPDH levels.
  - (b) by lowering plasma cholesterol and triglyceride levels, and the ratio of LDL to HDL cholesterol at doses which do not significantly modify cardiac GPDH levels.

The compounds of Formula 1 may therefore be used in therapy, in the treatment of conditions which can be alleviated by compounds which selectively mimic the effects of thyroid hormones in certain tissues whilst having little or no direct thyromimetic effect on the heart. For example, compounds of Formula 1 which raise hepatic GPDH levels and metabolic rate at doses which do not significantly modify cardiac GPDH levels are indicated in the treatment of obesity.

Agonists of Formula 1 will lower total plasma cholesterol, the ratio of LDL-cholesterol to HDL-cholesterol and triglyceride levels at doses which do not significantly modify cardiac GPDH levels are indicated for use as general antihyperlipidaemic (antihyperlipoproteinaemic) agents i.e. in the treatment of patients having elevated plasma lipid (cholesterol and triglyceride) levels. In

addition, in view of this effect on plasma cholesterol and triglyceride, they are also indicated for use as specific anti-hypercholesterolemic and anti-hypertriglyceridaemic agents.

Patients having elevated plasma lipid levels are considered at risk of developing coronary heart disease or other manifestations of atherosclerosis as a result of their high plasma cholesterol and/or triglyceride concentrations. Further, since LDL-cholesterol is believed to be the lipoprotein which induces atherosclerosis, and HDL-cholesterol believed to transport cholesterol from blood vessel walls to the liver and to prevent the build up of atherosclerotic plaque, anti-hyperlipidemic agents which lower the ratio of LDL-cholesterol to HDL cholesterol are indicated as anti-atherosclerotic agents, herein incorporated by reference U.S. patents 4,826,876 and 5,466,861.

The present invention also provides a method of producing selective thyromimetic activity in certain tissues except the heart which comprises administering to an animal in need thereof an effective amount to produce said activity of a compound of Formula 1 or a pharmaceutically acceptable salt thereof.

The present invention also relates to a method of lowering plasma lipid levels and a method of lowering the ratio of LDL-cholesterol to HDL-cholesterol levels by suitably administering a compound of this invention or a pharmaceutically acceptable 20 sale thereof.

In addition, compounds of Formula 1 may be indicated in thyroid hormone replacement therapy in patients with compromised cardiac function.

In therapeutic use the compounds of the present invention are usually administered in a standard pharmaceutical composition.

The present invention therefore provides in a further aspect pharmaceutical compositions comprising a compound of Formula 1 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier. Such compositions include those suitable for oral, parenteral or rectal administration.

# PHARMACEUTICAL COMPOSITIONS

Compounds of Formula 1 and their pharmaceutically acceptable salts which are active when given orally can be formulated as liquids for example syrups, 5 suspensions or emulsions, tablets, capsules and lozenges.

A liquid composition will generally consist of a suspension or solution of the compound or pharmaceutically acceptable salt in a suitable liquid carrier(s), for example ethanol, glycerine, sorbitol, non-aqueous solvent such as polyethylene glycol, oils or water, with a suspending agent, preservative, surfactant, wetting agent, 10 flavoring or coloring agent. Alternatively, a liquid formulation can be prepared from a reconstitutable powder.

For example a powder containing active compound, suspending agent, sucrose and a sweetener can be reconstituted with water to form a suspension; and a syrup can be prepared from a powder containing active ingredient, sucrose and a sweetener.

A composition in the form of a tablet can be prepared using any suitable pharmaceutical carrier(s) routinely used for preparing solid compositions. Examples of such carriers include magnesium stearate, starch, lactose, sucrose, microcrystalline cellulose and binders, for example polyvinylpyrrolidone. The tablet can also be provided with a color film coating, or color included as part of the carrier(s). In 20 addition, active compound can be formulated in a controlled release dosage form as a tablet comprising a hydrophilic or hydrophobic matrix.

A composition in the form of a capsule can be prepared using routine encapsulation procedures, for example by incorporation of active compound and excipients into a hard gelatin capsule. Alternatively, a semi-solid matrix of active compound and high molecular weight polyethylene glycol can be prepared and filled into a hard gelatin capsule; or a solution of active compound in polyethylene glycol or a suspension in edible oil, for example liquid paraffin or fractionated coconut oil can be prepared and filled into a soft gelatin capsule. Compound of Formula 1 and their pharmaceutically acceptable salts which are active when given parenterally can be 30 formulated for intramuscular or intravenous administration.

A typical composition for intra-muscular administration will consist of a suspension or solution of active ingredient in an oil, for example arachis oil or sesame oil. A typical composition for intravenous administration will consist of a sterile isotonic aqueous solution containing, for example active ingredient, dextrose, sodium

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WO 99/26966 PCT/US98/25296

chloride, a co-solvent, for example polyethylene glycol and, optionally, a chelating agent, for example ethylenediamine tetracetic acid and an anti-oxidant, for example, sodium metabisulphite. Alternatively, the solution can be freeze dried and then reconstituted with a suitable solvent just prior to administration.

Compounds of structure (1) and their pharmaceutically acceptable salts which are active on rectal administration can be formulated as suppositories. A typical suppository formulation will generally consist of active ingredient with a binding and/or lubricating agent such as a gelatin or cocoa butter or other low melting vegetable or synthetic wax or fat.

Compounds of Formula 1 and their pharmaceutically acceptable salts which are active on topical administration can be formulated as transdermal compositions. Such compositions include, for example, a backing, active compound reservoir, a control membrane, liner and contact adhesive.

The typical daily dose of a compound of Formula 1 varies according to 15 individual needs, the condition to be treated and with the route of administration. Suitable doses are in the general range of from 0.001 to 10 mg/kg bodyweight of the recipient per day.

Within this general dosage range, doses can be chosen at which the compounds of Formula 1 lower plasma cholesterol levels and raise metabolic rate 20 with little or no direct effect on the heart. In general, but not exclusively, such doses will be in the range of from lower doese (0.001 to 0.5 mg/kg) to higher doses (0.5 to 10 mg/kg).

In addition, within the general dose range, doses can be chosen at which the compounds of Formula 1 lower plasma cholesterol levels and have little or no effect 25 on the heart without raising metabolic rate. In general, but not exclusively, such doses will be in the range of from 0.001 to 0.5 mg/kg.

It is to be understood that the 2 sub ranges noted above are not mutually exclusive and that the particular activity encountered at a particular dose will depend on the nature of the compound of Formula 1 used.

Preferably, the compound of Formula 1 is in unit dosage form, for example, a tablet or a capsule so that the patient may self-administer a single dose. In general, unit doses contain in the range of from 0.05-100 mg of a compound of Formula 1. Preferred unit doses contain from 0.05 to 10 mg of a compound of Formula 1.

The active ingredient may be administered from 1 to 6 times a day. Thus daily doses are in general in the range of from 0.05 to 600 mg per day. Preferably, daily doses are in the range of from 0.05 to 100 mg per day. Most preferably from 0.05 to 5 mg per day.

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# <u>EXAMPLES</u>

#### **EXAMPLE 1 - SYNTHESIS OF TR LIGANDS**

Many TR ligands are known in the art, including T4 (thyroxine), T3, T2 and TS-9. See Jorgensen, Thyroid Hormones and Analogs, in *Hormonal Proteins and* 10 *Peptides, Thyroid Hormones* 107-204 (Choh Hao Li ed., 1978), incorporated by reference herein.

The syntheses of several TR ligands are described below.

# Synthesis of TS1, TS2, TS3, TS4, TS5

TS1, TS2, TS3, TS4 and TS5 and analogs thereof can all be prepared by simple acylation of the nitrogen atom of any thyronine analog, including T3 (3,5,3'-triiodo-L-thyronine), T4 (thyroxine) and 3,5-diiodothyronine. TS1 and TS2 are synthesized by reacting T3 with Ph<sub>2</sub>CHCO<sub>2</sub>NHS (N-hydroxy succinimide-2,2-diphenylacetate) and C<sub>16</sub>H<sub>33</sub>CO<sub>2</sub>NHS, respectively. TS3 is synthesized by reacting T3 with FMOC-Cl (fluorenylmethyloxycarbonylchloride). TS4 is synthesized by reacting T3 with tBOC<sub>2</sub>O (tBOC anhydride or di-t-butyldicarbonate). TS5, which differs from TS1-4 by having a -H instead of an -I at the R'<sub>3</sub> position, is synthesized by reacting 3,5-diiodothyronine with tBOC<sub>2</sub>O. The general reaction scheme for TS1, TS2, TS3, TS4 and TS5 is depicted in FIG. 11. It should be noted that in the reaction scheme, both TS5 and its precursor both have a hydrogen rather than an iodine at the R'<sub>3</sub> position.

# Synthesis of TS6 and TS7

TS6 is synthesized by reacting TS5 with paranitrophenylisocyanate. TS7 is synthesized by reacting TS6 with TFA (trifluoroacetic acid), which cleaves the tBOC group. These reactions are simple organic synthesis reactions that can be performed by anyone of ordinary skill in the art. The synthetic scheme for TS6 and TS7 is diagrammed in **FIG. 12**.

# Synthesis of TS8

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TS8 is synthesized by reacting TS5 with Ph<sub>2</sub>CHNH<sub>2</sub> (diphenylmethylamine) in the presence of triethylamine and any amide forming condensing reagent, such as 5 TBTU (hydroxybenztriazoleuronium tetrafluoroborate) or HBTU (hydroxybenztriazoleuronium hexafluorophosphate). The synthesis scheme for TS8 is depicted in FIG. 13.

# SYNTHESIS OF 3,5-DIIODO-3'ISOPROPYLTHYRONINE DERIVATIVES

For designing a class of antagonists, it is important to have a hydrophobic group at the 3' position as well as an extension at the 5' position. Preferred hydrophobic groups at the 3' position include: methyl, benzyl, phenyl, iodo, and heterocyclic structures. The synthesis of a 3,5-diiodo-3'-isopropyl-5'-substituted thyronine is described below. The example provided describes the specific steps for synthesizing the TS10 compound, but this general reaction scheme can be used by one of ordinary skill in the art to synthesize any number of 3,5,-diiodo-3'-isopropyl-5'-substituted thyronine derivatives, which are characterized by having an extension at the 5' position. Additional compounds of this class can be synthesized using known organic synthesis techniques.

The synthesis of TS10 is described below and is depicted in **FIG. 14.** Numbers used in the reaction scheme for TS10 indicating the reaction product for each step are in parentheses.

2-Formyl-6-isopropylanisole (1): 2-formyl-6-isopropylanisole (10.0 g, 61 mmol), as made by Casiraghi, *et al.* JCS Perkin I, 1862 (1980) (incorporated by 25 reference), is added dropwise to a suspension of sodium hydride (3.7 g, 153 mmol) in 50 mL THF and 50 mL of DMF in a round bottom flask. The addition generates an exothermic reaction and formation of a gray solid. Methyl iodide (26.0 g, 183 mmol) is then added dropwise and the reaction mixture is stirred at room temperature for 5 hours. The reaction mixture is quenched with 20 mL of water, then poured into 500 mL of water, and is extracted with ether (2 x 300 mL). The ether layers are combined, washed with water (5 x 1000 mL), dried over magnesium sulfate and concentrated in vacuo to provide 10.2 g (94%) of the title compound, with the following <sup>1</sup>H NMR (CDCl<sub>3</sub>) properties: d 10.30 (s, 1H), 7.63 (d, 1H, J=3 Hz), 7.50

(d, 1H, J=3 Hz), 7.13 (t, 1H, J=3 Hz), 3.81 (s, 3H), 3.31 (heptet, 1H, J=7.5 Hz), 1.19 (d, 6H, J=7.5 Hz).

2-(2-Hydroxynonyl)-6-isopropylanisole (not shown in scheme):

5 Octylmagnesium chloride (8.4 mL, 16.9 mmol, 2.0 M) is added dropwise to a solution of 1 (1.5 g, 8.4 mmol) in 10 mL THF at -78°C. The reaction mixture is stirred for 2 hours with warming to room temperature. The reaction mixture is diluted with 50 mL ether and poured into 50 mL water. The ether layer is washed with brine (1 x 50 mL), dried over sodium sulfate, and concentrated in vacuo. Flash chromatography (silica 10 gel, 10% ether/hexane → 15% ether/hexane) provides 734 mg (30%) of the title compound with the following <sup>1</sup>H NMR (CDCl<sub>3</sub>) properties: d 7.33-7.10 (m, 3H), 5.00 (br. s, 1H), 3.81 (s, 3H), 3.33 (heptet, 1H, J=7 Hz) 1.90-1.19 (m, 14H), 0.86 (t, 3H, J=6.5 Hz); HRMS (EI), found: 292.2404; calc'd: 292.2402.

2-nonyl-6-isopropylanisole (2): Compound 2 (663 mg, 2.3 mmol) is dissolved in solution of 5 mL ethanol and 5 mL acetic acid, and a spatula tip of palladium on carbon catalyst is added. The reaction mixture is then charged with hydrogen gas (using a simple balloon and needle) and the mixture is stirred at room temperature overnight. The next day, the reaction mixture is poured into ether (100 mL) and the ether layer is extracted with saturated sodium bicarbonate (3 x 100 mL). The ether layer is dried over sodium sulfate and concentrated *in vacuo* to provide 581 mg (91%) of (2) with the following <sup>1</sup>H NMR (CDCl<sub>3</sub>) properties: d 7.14-7.00 (m, 3H), 3.75 (s, 3H), 3.36 (heptet, 1H, J=6.8 Hz), 2.63 (t, 2H, J=7.5 Hz), 1.68-1.15 (m, 14H), 0.86 (t, 3H, J=5.5 Hz); HRMS (EI), mass found: 276.2459; calculated: 276.2453.

Thyronine adduct (4): Fuming nitric acid (0.071 mL) is added to 0.184 mL acetic anhydride chilled to -5°C. Iodine (66 mg) is added to this mixture followed by trifluoroacetic acid (0.124 mL). This mixture is stirred for 1 hour with warming to room temperature, at which point all of the iodine is dissolved. The reaction mixture was then concentrated *in vacuo* to provide an oily semi-solid material. The residue was dissolved in 0.7 mL of acetic anhydride and cooled to -20°C. A solution of anisole (2) (581 mg, 2.1 mmol) in 1.2 mL acetic anhydride and 0.58 mL TFA is added dropwise. The reaction mixture is stirred at -20° for 1 hour, then stirred overnight with warming to room temperature. The reaction mixture is partitioned between water and methylene chloride. The methylene chloride layer is dried over sodium

sulfate and concentrated *in vacuo* to provide the iodonium salt (3) as an oil. This material is not purified or characterized, and is directly introduced into the coupling reaction.

N-Trifluoroacetyl-3,5-diiodotyrosine methyl ester (552 mg, 1.0 mmol) prepared according to the procedure of N. Lewis and P. Wallbank, Synthesis 1103 (1987) (incorporated by reference) and all of the crude iodonium salt (3) from above is dissolved in 5 mL of anhydrous methanol. Diazabicyclo[5.4.0]undecane (DBU) (183 mg, 1.2 mmol) and a spatula tip of copper-bronze are added and the resulting mixture is stirred at room temperature overnight. The next day, the reaction mixture is filtered, and the filtrate is concentrated in vacuo. The crude residue is purified by flash chromatography (silica gel, 10% ethyl acetate/hexane) to provide 30 mg (4%) of the protected thyronine adduct (4).

Deprotected thyronine (TS10): The protected thyronine 4 (30 mg, 0.04 mmol) is dissolved in a mixture of 2.25 mL acetic acid and 2.25 mL 49% hydrobromic acid.

15 The reaction mixture is heated to reflux for 5 hours. The reaction mixture is cooled to room temperature, and the solvents are removed *in vacuo*. Water is added to triturate the oily residue into a gray solid. This solid material is filtered, washed with water, and dried over P<sub>2</sub>O<sub>5</sub> *in vacuo* to provide 24 mg (81%) of the title compound, TS10, with the following <sup>1</sup>H NMR (CDCl<sub>3</sub>) properties: d 7.57 (s, 1H), 6.86 (s, 1H), 6.45 (s, 20 1H), 6.34 (s, 1H), 4.81 (m, 1H), 3.86 (s, 3H), 3.71 (s, 3H), 3.33-3.05 (m, 3H), 2.58-2.47 (m, 2H), 1.62-0.76 (m, 23H); MS (LSIMS): M<sup>+</sup> = 817.0.

As mentioned above, this reaction scheme can be modified by one of ordinary skill in the art to synthesize a class of compounds characterized by 3,5-diiodo-3'isopropylthyronine derivatives, wherein (1) the 3' isopropyl group can be replaced with a hydrophobic group, including methyl, benzyl, phenyl, iodo, and heterocyclic structures, and (2) a wide variety of chemical structures can be incorporated at the 5' position, including alkyl groups, planar aryl, heterocyclic groups, or polar and/or charged groups.

The aldehyde (1) in the above reaction scheme is a versatile synthetic 30 intermediate which allows for the attachment of a variety of chemical moieties to the 5' position of the final thyronine derivative. In addition, a variety of chemical reactions can be used to attach the chemical moieties. These reactions are well known in the art and include organometallic additions to the aldehyde (including Grignard reagents, organolithiums, etc.), reductive amination reactions of the aldehyde with a

primary or secondary amine, and Wittig olefination reactions with a phosphorous ylid or stabilized phosphonate anion. Other possibilities include reduction of the aldehyde to a benzyl alcohol allowing for etherification reactions at the 5' position. As mentioned above, these methods allow for a wide variety of chemical structures to be incorporated at the 5' position of the final thyronine derivative, including alkyl groups, planar aryl, heterocyclic groups or polar and/or charged groups.

Synthesis of 3, 5-dibromo-4-(3',5'-diisopropyl-4'-hydroxyphenoxy) benzoic acid (Compound 11).

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- (a) A mixture of 2,6-diisopropyl phenol (20 g, 0.11 mol), potassium carbonate (62 g, 0.45 mol), acetone (160 ml) and methyl iodide (28 ml, 0.45 mole) is refluxed for three days. The reaction mixture is filtered through celite, evaporated, dissolved in ether, washed twice with 1M sodium hydroxide, dried over magnesium sulphate and concentrated to afford 15.1 g (0.08 mol, 70%) of 2,6-diisopropyl anisole as a slightly yellow oil.
- (b) Fuming nitric acid (12.4 ml, 265 mmol) is added dropwise to 31.4 ml of acetic anhydride which is cooled in a dry ice/carbon tetrachloride bath. Iodine 11.3 g, 44.4 mmol) is added in one portion followed by dropwise addition of trifluoroacetic acid (20.5 ml, 266 mmole). The reaction mixture is stirred at room temperature until all the iodine is dissolved. Nitrogen oxides are removed by flushing nitrogen into the vessel. The reaction mixture is concentrated, the residue is dissolved in 126 ml of acetic anhydride and is cooled in a dry ice/carbon tetrachloride bath. To the stirred solution 2,6-diisopropylanisole (51 g, 266 mmol) in 150 ml of acetic anhydride and 22.6 ml of trifluoroacetic acid is added dropwise. The reaction mixture is left to stand at room temperature over night and then is concentrated. The residue is taken up in 150 ml of methanol and treated with 150 ml of 10% aqueous sodium bisulfite solution and 1 liter of 2M sodium borotetrafluoride solution. After the precipitate aggregates, petroleum ether is added and the supernatant is decanted. The precipitate is triturated with petroleum ether, filtered, washed with petroleum ether and dried at room

temperature in vacuo. This affords 34 g (57 mmol, 65%) of bis(3,5-diisopropyl-4-methoxyphenyl)iodonium tetrafluoroborate as a white solid.

- (c) To a stirred solution of 3,5-dibromo-4-hydroxybenzoic acid (12 g, 40.5 mmol) in 250 ml of methanol, thionyl chloride (3 ml) is added dropwise. The 5 reaction mixture is refluxed for five days, water is added and the precipitated product is filtered off. The residue is dissolved in ethyl acetate. From the aqueous phase, methanol is removed by concentration. The aqueous phase is then saturated with sodium chloride, and extracted with ethyl acetate. The combined organic phases are dried over magnesium sulphate, filtered and concentrated. This gives 12.5 g (40.5 mmol, 100%) of 3,5-dibromo-4-hydroxymethyl benzoate as a white crystalline solid.
- (d) The products obtained in steps b and c are reacted with each other according to the following protocol. To bis(3,5-diisopropyl-4-methoxyphenyl)iodonium tetrafluoroborate (2.86 g, 4.8 mmole) and copper bronze (0.42 g, 6.4 mmole) in 7 ml. of dichloromethane at 0°C is added dropwise a solution of 3,5-dibromo-4-hydroxymethyl benzoate (1.0 g, 3.2 mmole) and triethylamine (0.36 g, 3.5 mmole) in 5 ml of dichloromethane. The reaction mixture is stirred in the dark for eight days and then is filtered through celite. The filtrate is concentrated and the residue is purified by column chromatography (silica gel, 97:3 petroleum ether/ethyl acetate) to give 0.62 g (1.2 mmole, 39%) of 3,5-dibromo-4-(3',5'-diisopropyl-4'-20 methoxyphenoxy)methyl benzoate as a solid.
- The product from step d (0.2 g, 0.4 mmole) is dissolved in 2 ml. dichloromethane, is put under nitrogen and is cooled at -40°C. To the stirred solution is added 1M BBr<sub>3</sub> (1.2 ml, 1.2 mmole) dropwise. The reaction mixture is allowed to reach room temperature and then is left over night. It is cooled to 0°C and then 25 hydrolyzed with water. Dichloromethane is removed by concentration and the aqueous phase is extracted with ethyl acetate. The organic phase is washed with 1M hydrochloric acid and brine. Then it is dried over magnesium sulphate, filtered and concentrated. The residue is chromatographed (silica, 96:3.6:0.4 dichloromethane/methanol/acetic acid) producing 93 mg (0.2 mmole, 51%) of 3,5-30 dibromo-4-(3',5'-diisopropyl-4'-hydroxyphenoxy)benzoic acid as a white solid. <sup>1</sup>H nmr (CDCl<sub>3</sub>) δ 1.23 (d, 12H, methyl), 3.11 (m, 2H, CH), 6.50 (s, 2H, 2,6-H) 8.33 (s, 2H, 2',6'-H).

Synthesis of addition ligands are described in U.S. Serial No. 08/877,792, filed June 18, 1997 which is herein incorporated in its entirety by reference.

**TABLE 2** and **FIG. 15** depict the structures of several TR ligands in reference to Formula I.

5

TABLE 2

Cmpd	R <sub>3</sub>	R4	R <sub>5</sub>	R'3	R' <sub>4</sub>	R's	R <sub>1</sub>
*T <sub>3</sub>	-I	-0-	-I	-I	-ОН	-H	-CH₂CH(NH₂)CO₂H
*T <sub>4</sub>	-I	<b>-</b> O-	-I	-1	-OH	-I	-ĊH₂CH(NH₂)CO₂H
TS1	-l	-0-	-I	-l	-OH	-Н	-CH₂CH[NHCOCH∳₂]CO₂H
TS2	-I	-0-	-1	-I	-ОН	-Н	-CH <sub>2</sub> CH[NHCO(CH <sub>2</sub> ) <sub>15</sub> CH <sub>3</sub> ]CO <sub>2</sub> H
TS3	-I	-O-	-1	-1	-OH	-Н	-CH₂CH[NH-FMOC]CO₂H
TS4	-J	-0-	-1	-I	-OH	-Н	-CH₂CH[NH-tBOC]CO₂H
TS5	-1	-0-	-1	-H	-ОН	-Н	-CH₂CH[NH-1BOC]CO₂H
TS6	-J	-0-	-I	-H	-OC(O)NH=Ø <sub>p</sub> NO <sub>2</sub>	-H	-CH₂CH[NH-tBOC]CO₂H
TS7	-I	<b>-</b> O-	-I	-i	- OC(O)NH=NHØNO <sub>2</sub>	-H	-CH₂CH(NH₂)CO₂H
TS8	I	-0-	-I	-Н	-NH-CHØØ	-H	-CH <sub>2</sub> CH[NH-tBOC]CO <sub>2</sub> H
TS9	-I	-O-	-I	-IsoPr	-ОН	-Н	-CH₂CH(NH₂)CO₂H
TS10	-I	-0-	-I	-IsoPr	-ОН	-(CH) <sub>8</sub> - CH <sub>3</sub>	-CH₂CH(NH₂)CO₂H

\*

Prior Art Compound

-Ø:

phenyl

-ØpNO<sub>2</sub>:

para nitro phenyl

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# **EXAMPLE 2 - RECEPTOR BINDING ASSAYS OF TR LIGANDS**

To test the ability of synthesized TR ligands to bind to a thyroid receptor (TR), the binding affinity of a TR ligand for TR is assayed using TR's prepared from rat liver nuclei and [125]T<sub>3</sub> as described in J.D. Apriletti, J.B. Baxter, and T.N. Lavin, J. Biol. Chem., 263: 9409-9417 (1988). The apparent Kd's are calculated using the method described by Apriletti (1995) and Apriletti (1988). The apparent Kd's are presented in TABLE 3. The apparent Kd's (App.Kd) are determined in the presence

of the sample to be assayed, 1 nM [<sup>125</sup>I]T<sub>3</sub>, and 50Tg/ml core histones, in buffer E (400 mM KCl, 200 mM potassium phosphate, pH 8.0, 0.5 mM EDTA, 1 mM MgCl<sub>2</sub>, 10% glycerol, 1 mM DTT) in a volume of 0.21 ml. After incubation overnight at 4°C, 0.2 ml of the incubation mixture is loaded onto a Quick-Sep Sephadex G-25 column 5 (2.7 x 0.9 cm, 1.7 ml bed volume) equilibrated with buffer E. The excluded peak of protein-bound [<sup>125</sup>I]T<sub>3</sub> is eluted with 1 ml of buffer E, collected in a test tube, and counted. Specific T<sub>3</sub> binding is calculated by subtracting nonspecific binding from total binding.

TABLE 3

Compound	App.Kd(nM)	Coactivation Assay RIP-140	EC <sub>50</sub> (M)
T <sub>3</sub>	0.06	+	10-10
T <sub>4</sub>	2	+	10-9
TS1	4	+	10-7
TS2	1400	nd	nd
TS3	4	+ .	10-8
TS4	8	+	nd
TS5	220	+	10-6
TS6 .	>10000	nd	nd
TS7	260	+	10-7
TS8	6000	nd	nd
TS9	1	+	10 <sup>-10</sup>
TS10	. 400	+	10-6

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+: RIP-140 Binding

-: RIP-140 Binding

nd: Not Determined

# **EXAMPLE 3 - INCREASED NUCLEAR PROTEIN COACTIVATION BY TR LIGANDS**

To test the ability of TR ligands to activate the binding of TR to the nuclear activation protein RIP-140 (a nuclear protein that can bind to nuclear receptors, such as the estrogen receptor), a TR ligand is liganded to TR and then incubated with RIP-140 as described in V. Cavailles, et al., EMBO J., 14(15):3741-3751 (1995), which is incorporated by reference herein. In this assay, <sup>35</sup>S-RIP-140 protein binds to liganded TR but not unliganded TR. Many TR <sup>35</sup>S ligands can activate RIP-140 binding as shown in **TABLE 3**.

# 10 Example 4 - TR LIGAND BINDING AND TR ACTIVATION IN CULTURED CELLS

To test TR activation of transcription in a cellular environment, TR ligands are assayed for their ability to activate a reporter gene, chloramphenicol transferase ("CAT"), which has a TR DNA binding sequence operatively linked to it. Either GC or L937 cells (available from the ATCC) can be used, respectively). In such assays, a 15 TR ligand crosses the cell membrane, binds to the TR, and activates the TR, which in turn activates gene transcription of the CAT by binding the TR DNA binding region upstream of the CAT gene. The effective concentration for half maximal gene activation (EC50) is determined by assaying CAT gene activation at various concentrations as described herein and in the literature. The results of CAT gene 20 activation experiments are shown in TABLE 3.

# **CAT GENE ACTIVATION ASSAYS**

Functional response to thyroid hormone (3,5,3'-triiodo-L-thyronine, T<sub>3</sub>) and TR ligands is assessed either in a rat pituitary cell line, GC cells, that contain endogenous thyroid hormone receptors (TRs) or U937 cells that contain exogenous TRs expressed as known in the art. GC cells are grown in 10-cm dishes in RPMI 1640 with 10% newborn bovine serum, 2 mM glutamine, 50 units/ml penicillin and 50 Tg/ml streptomycin. For transfections, cells are trypsinized, resuspended in buffer (PBS, 0.1% glucose) and mixed with a TREtkCAT plasmid (10 mg) or phage in 0.5 ml buffer (15±5 million cells) and electroporated using a Bio-Rad gene pulser at 0.33 kvolts and 960 mF. The TREtkCAT plasmid contains two copies of a T<sub>3</sub> response element (AGGTCAcaggAGGTCA) cloned in the Hind III site of the pUC19 polylinker immediately upstream of a minimal (-32/+45) thymidine kinase promoter

linked to CAT (tkCAT) coding sequences. After electroporation, cells are pooled in growth medium (RPMI with 10% charcoal-treated, hormone stripped, newborn bovine serum), plated in 6-well dishes and treated with either ethanol or hormone. CAT activity is determined 24 hours later as described D. C. Leitman, R. C. J. 8 Ribeiro, E. R. Mackow, J. D. Baxter, B. L. West, *J. Biol. Chem.* 266, 9343 (1991), which is incorporated by reference herein.

EFFECT OF TS-10 ON THE TRANSCRIPTIONAL REGULATION OF THE DR4-ALP REPORTER GENE IN THE PRESENCE OR ABSENCE OF T3.

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Characteristics of the TRAF cells: TRAFa1 are CHO K1 cells stably transformed with an expression vector encoding the human thyroid hormone receptor α1 and a DR4,ALP reporter vector; TRAFb1 are CHO K1 cells stably transformed with an expression vector encoding the human thyroid hormone receptor β1 and a 15 DR4-ALP reporter vector.

Interpretation of the effect of compound TS-10 on the transcriptional regulation of the DR4-ALP reporter gene in the presence or absence of T3.

TRAFa1 reporter cells: TS-10 alone (open circles) induces a partial activation of the expression of the ALP reporter protein amounting to approximately 27% of the maximal effect by the natural thyroid hormone T3. In the presence of T3 (filled circles), TS-10 has a weak antagonistic effect. The EC50 concentration for the agonistic effect of TS-10 and the EC50 concentration for its T3 antagonistic effect, 25 respectively, is indicated in FIG. 18.

In FIG. 18, open and filled circles with dotted lines show the dose-dependent effect of TS-10/T3 on the toxicity marker (MTS/PMS), reduction of tetrazolium salt in the mitochondria, displayed on the right y-axis as optical density. There is no obvious toxic effect of TS-10 on the MTS-PMS marker but there is a clear effect on the morphology of the cells, as can be seen under the light microscope, at the highest concentration of TS-10 (32 mM) both in the absence and presence of T3, respectively (not shown in the figure).

TRAFb1 reporter cells: TS-10 alone (open circles) induces a partial activation of the expression of the ALP reporter protein amounting to approximately 35% of the maximal effect by T3. The EC50 concentration for the agonistic effect of 5 TS-10 is indicated in FIG. 19. In the presence of T3 (filled circles), TS-10 shows, if anything, a slight potentiation of the T3 effect on the expression of the ALP reporter protein. The T3 inhibitory effect of TS-10 at its highest concentration used (32 mM) is a toxic effect rather than T3 antagonism.

In FIG. 19, open and filled circles with dotted lines show the dose-dependent effect of TS-10/T3 on the toxicity marker (MTS/PMS), reduction of tetrazolium salt in the mitochondria, displayed on the right y-axis as optical density. There is no obvious toxic effect of TS-10 on the MTS-PMS marker but a clear effect on the morphology of the cells can be observed, under the light microscope, at the highest concentration of TS-10 (32 mM) both in the absence and presence of T3, respectively (not shown in the figure).

HepG2 (HAF18) reporter cells: TS-10 alone (open circles) induces a partial activation of the expression of the ALP reporter protein amounting to slightly more 20 than 50% of the maximal effect by T3. The EC50 concentration for the agonistic effect of TS-10 is indicated in FIG. 20. In the presence of T3 (filled circles), TS-10 shows no effect i.e. no T3 antagonism nor potentiation/additive effect to T3. Open and filled circles with dotted lines show the dose-dependent effect of TS-10/T3 on the toxicity marker (MTS/PMS), reduction of tetrazolium salt in the mitochondria, 25 displayed on the right y-axis as optical density. There is no obvious toxic effect of TS-10 on the MTS/PMS marker or on the morphology of the cells, as can be observed using a light microscope, at any concentration of TS-10/T3 used.

# Example 5 - Comparisons of Human TR-α and Human TR-β

# Competition for [125I]T<sub>3</sub> binding to TR LBD by T<sub>3</sub> and Triac

The drug, Triac, is a thyroid hormone agonist. Triac is 3,5,3'-triiodothyroacetic acid and is described in Jorgensen, Thyroid Hormones and Analogs in *Hormonal Proteins and Peptides, Thyroid Hormones* at 150-151 (1978). Another

5

WO 99/26966 PCT/US98/25296

compound that can be used in place of Triac is 3,5-diiodo-3'-isopropylthyroacetic acid. Competition assays are performed to compare the displacement of  $[^{125}I]T_3$  from binding with human TR- $\alpha$  LBD or human TR- $\beta$  LBD by unlabeled  $T_3$  or Triac. The results of such assays are depicted in FIG. 16.

Standard binding reactions are prepared containing 1 nM [ $^{125}I$ ]T<sub>3</sub>, 30 fmol of human TR- $\alpha$  (empty symbols) or  $\beta$  (solid symbols), and various concentrations of competing unlabeled T<sub>3</sub> (circles) or Triac (triangles). Assays are performed in duplicate.

# Competition for [125I]T<sub>3</sub> binding to variant TR LBD by T<sub>3</sub>, Triac and GC-1

The following assays residues involved in selective binding among TR isoforms. Competition assays are performed to compare the displacement of [125]T<sub>3</sub> from binding with wild-type human TR-α LBD or human TR-β LBD, to a variant form of the TR LBDs by unlabeled T<sub>3</sub>, Triac or GC-1. A variant TR-α or TR-β is constructed by substituting an amino acid found in the corresponding position of the other TR isoform. For example, asparagine 331 in human TR-β corresponds to serine 277 in human TR-α. To test binding specificity contributed by this position, a variant human TR-β is constructed that contains asparagine 331 substituted with a serine residue (designated Asn331Ser or N331S). Binding assays are described in *Apriletti* 20 et al. (Protein Expression and Purification 6:363-370 (1995)). The results of such assays are depicted in FIG. 27, and summarized in Table 4 below.

TABLE 4
Effect of TR-β Substitution N331S on Binding Affinity

Ligand	Native TR-α	Native TR-β	Mutant TR-β
Т3	20 pM	60 pM	100 pM
T4	600	3000	ND
Triac	20	20	100
IpBr <sub>2</sub>	17	ND	ND
Dimit	6000	8000	ND
GC-1	200	40	400

Competition curves comparing wildtype TR-β versus the variant TR-β N331S for 5 binding T3, Triac or GC-1 show that the affinity of the mutant receptor for Triac was reduced to approximately the same as for T3 (vs. 3-fold greater in wild type) so that the relative affinities are similar to wild-type TR-α. The affinity for GC-1 was also reduced to several fold less than T3, as is seen with TR-α.

Comparison of the affinity of TR- $\beta$  variant N331S to the native TRs for 10 selected ligands is as follows:

Native TR- $\alpha$  for various ligands (T3, T4, Triac, IpBr2, Dimit, GC-1): IpBr<sub>2</sub>. > Triac  $\cong$  T3 > GC-1 > T4 > Dimit

Native TR-β (T3, T4, Triac, Dimit, GC-1)

Triac > GC-1  $\geq$  T3 > T4 > Dimit

15 Variant TR-β (N331S) (T3, Triac, GC-1)

Triac  $\cong$  T3 > GC-1.

# Scatchard Analysis of [125I]T<sub>3</sub> Binding to TR

Human TR- $\alpha$  (left panel) or human TR- $\beta$  (right panel) is assayed for T<sub>3</sub> 20 binding in the presence of increasing concentrations of [ $^{125}I$ ]T<sub>3</sub>. The apparent equilibrium dissociation constant (20 pM for I and 67 pM for  $\beta$ ) is calculated by linear regression analysis and is depicted in **FIG. 17**.

3, 5-DIBROMO-4-(3',5'-DIISOPROPYL-4'-HYDROXYPHENOXY) BENZOIC ACID IS A TR-α SELECTIVE SYNTHETIC LIGAND.

3, 5-dibromo-4-(3',5'-diisopropyl-4'-hydroxyphenoxy) benzoic acid (Compound 11), the structure of which is drawn above, is assayed for binding to the two different isoforms of the TR, TR-α and TR-β. Compound 11 exhibits an IC50 of 1.6 TM for binding to TR-α and an IC50 of 0.91 TM for binding to TR-β. Assays for determining selective binding to the TR-α or TR-β LBD can include reporter assays, 15 as described herein. See also Hollenberg, et al., J. Biol. Chem., (1995) 270(24):14274-14280.

#### Example 6 - Preparation and Purification of a $TR-\alpha$ LBD

Rat TR-α LBD, residues Met122 - Val410, is purified from *E. coli* ("LBD-20 122/410"). The expression vector encoding the rat TR-α LBD is freshly transfected into *E. coli* strain BL21(DE3) and grown at 22°C in a 50-liter fermenter using 2x LB medium. At an A<sub>600</sub> of 2.5-3, IPTG is added to 0.5 mM and growth is continued for 3 h before harvesting. The bacterial pellet is quickly frozen in liquid nitrogen and stored at -70°C until processed. Extraction and purification steps are carried out at 25 4°C. The bacteria are thawed in extraction buffer (20MM Hepes, pH 8.-, 1 mM EDTA, 0.1% MTG, 0.1 mM PMSF, and 10% glycerol) at a ratio of 10 ml buffer/g bacteria. Bacteria are lysed by incubation for 15 min. with 0.2 mg/ml lysozyme and sonicated at maximum power while simultaneously homogenized with a Brinkmann homogenizer (Model PT 10/35 with generator PTA 35/2) until the solution loses its viscosity. After centrifugation for 10 min at 10,000 g, the supernatant is adjusted to 0.4 M KCl, treated with 0.6% PEI to precipitate fragmented DNA, and centrifuged for 10 min at 10,000 g. The rat TR-α LBD in the supernatant is then precipitated with 50% ammonium sulfate and centrifuged for 10 min at 10,000 g. The precipitate is

5

Page 76 of 447

PCT/US98/25296 WO 99/26966

resuspended with buffer B (20 mM Hepes, pH 8.0, 1 mM EDTA, 1 mM DTT, 0.1 mM PMSF, 0.01% Lubrol, and 10% glycerol) to a final conductivity of 9 mS/cm (approx. 0.7 M ammonium sulfate) and centrifuged 1 h at 100,000g. The supernatant is frozen in liquid nitrogen and stored at -70°C.

The crude extract is thawed, bound with a tracer amount of [125I]T3, and loaded directly onto a phenyl-Toyopearl hydrophobic interaction column (2.6 x 18 cm, 95 ml bed volume) at 1.5 ml/min. The column is eluted with a 2-h gradient from 0.7 ammonium sulfate, no glycerol to no salt, 20% glycerol in buffer C (20 mM Hepes, pH 8.0, 0.5 mM EDTA, 1 mM DTT, 0.2 mM PMSF). The rat TR-α LBD 10 prebound to tracer [125I]T<sub>3</sub> (less than 0.005% of total rat TR-α LBD) is detected using a flow-through gamma emission detector, whereas unliganded rat TR-I LBD is assayed by postcolumn [125I]T<sub>3</sub> binding assays (described herein).

The phenyl-Toyopearl unliganded rat TR-a LBD peak fractions are pooled, diluted with buffer B to a conductivity of 0.5 mS/cm (equivalent to approx. 20 mM 15 ammonium sulfate), loaded onto a TSK-DEAE anion-exchange column (2 x 15 cm, 47 ml bed volume) at 4 ml/min, and eluted with a 60-min gradient from 50 to 200 mM NaCl in buffer B.

The unliganded rat TR-\alpha LBD peak fractions from TSK-DEAE are pooled, diluted twofold with buffer B, loaded at 0.75 ml/min on a TSK-heparin HPLC column 20 (0.8 x 7.5 cm, 3 ml bed volume), and eluted with a 50 to 400 mM NaCl gradient in buffer B.

The pool of unliganded rat TR-α LBD peak fractions from the TSK-heparin column is adjusted to 0.7 M ammonium sulfate, loaded at 0.75 ml/min on a TSKphenyl HPLC column (0.8 x 7.5 cm, 3 ml bed volume), and eluted with a 60-min 25 gradient from 0.7 M ammonium sulfate without glycerol to no salt with 20% glycerol in buffer C. The fractions containing unliganded rat TR-α LBD are pooled and incubated with a five fold excess of hormone for 1 h, the salt concentration is adjusted to 0.7 M ammonium sulfate, and the sample is reloaded and chromatographed on the same column as described above.

#### Example 7 - Crystallization of Liganded TR-α LBD

Material from a single LBD-122/410 preparation is divided into batches, and quantitatively bound with one of the following ligands: Dimit, T<sub>3</sub>, or Triac IpBr<sub>2</sub> (3,5dibromo-3'isopropylthyronine) for the final purification step.

To maintain full saturation of rat TR-α LBD with a ligand, and to prepare the complex for crystallization, the ligand-bound rat TR-α LBD is concentrated and desalted in an Amicon Centricon-10 microconcentrator (McGrath et al, *Biotechniques*, (1989) 7:246-247, incorporated by reference herein), using 10 mM Hepes (pH 7.0), 3.0 mM DTT, and 1.0 nM to 10 nM ligand.

10 Factorial crystallization screening trials (Jancarik & Kim, J. Appl. Crystallogr. (1991) 24:409-411, incorporated by reference herein) are carried out for rat TR-α LBD bound to selected ligands using hanging-drop vapor diffusion at 17°C (with 1 µl protein solution, 1 µl precipitant solution and a 0.5 ml reservoir using silanized coverslip: (McPherson, Preparation and Analysis of Protein Crystals (1982), 15 incorporated by reference herein). Rat  $TR-\alpha$  LBD is not stable at 4°C and is stored at -80°C, where it maintains its avidity for hormone and its crystallizability for approximately two to three months. These procedures are carried out as described in McGrath, M.E. et al., J. Mol. Biol. (1994) 237:236-239 (incorporated by reference). Crystals are obtained in condition 21 of the screening trials (Jancarik & Kim 1991) 20 and conditions are then optimized. Wedge-shaped crystals are reproducibly obtained with hanging-drop vapor fusion at 22°C with 15% 2-methyl-2,4-pentanediol (MPD), 0.2 M ammonium acetate and 0.1 M sodium cacodylate (pH 6.7), 3 mM DTT, with 2 μl protein solution, 1 μl precipitant solution and a 0.6 ml reservoir using silanized coverslip, and with 8.7 mg/ml (Dimit), 5.5 mg/ml (IpBr<sub>2</sub>), 5 mg/ml (Triac), or 2.3 25 mg/ml (T<sub>3</sub>) over a period of three days. Under these conditions, diffraction quality crystals (dimension 0.5 x 0.2 x 0.0075 mm<sup>3</sup>) can be grown at ambient temperature (22°C). The best crystals have a limiting dimension of approximately 100 Tm and are obtained at a protein concentration between 2.3 and 8.7 mg/ml in the presence of 3 mM DTT. The crystals are of the monoclinic space group C2, with one monomer in 30 the asymmetric unit.

# Example 8 - Crystallization of Human TR- $\beta$ LBD Complexed with T3, Triac, or GC-1

Human TR-β LBD complexed with T<sub>3</sub>, Triac, or GC-1 are purified according to the same procedures described above for the rat TR-α LBD, with the following 5 modifications.

The expression of human TR-β LBD differs from the rat TR-α LBD in that the human TR-β LBD residues extend from the amino acid at position 716 through the amino acid at position 1022, according to the amino acid numbering scheme for the various nuclear receptor LBDs depicted in FIG. 3. FIG. 3 illustrates a numbering scheme applicable to all of the nuclear receptors listed as well as to any additional homologous nuclear receptors. The vertical lines on FIG. 3 at position 725 and at position 1025 delineate the preferred minimum amino acid sequence necessary to obtain adequate binding of ligand. The amino acid sequence from position 716 to position 1022 according to the numbering scheme of FIG. 3 corresponds to the amino 15 acid positions 202 to 461 according to the conventional numbering of the amino acid sequence of human TR-β which is publicly available. Also, the human TR-β LBD is expressed with a histidine tag, as described in Crowe *et al.*, *Methods in Molecular Biology* (1994) 31:371-387, incorporated by reference herein.

The purification of human TR-β LBD is the same as that described above for 20 the rat TR-α LBD with the following exceptions. First, before the purification step using the hydrophobic interaction column, a step is added in which the expressed human TR-β LBD is purified using a nickel NTA column (commercially available from Qiagen, Chatsworth, CA) according to manufacturer's instructions, and eluted with 200 mM imidazole. The second difference is that in the purification of the 25 human TR-β LBD, the purification step using a heparin column is omitted.

The crystallization of human TR-β LBD bound to T<sub>3</sub>, Triac or GC-1 is as follows. Crystals are obtained in condition 7 of the factorial screen using hanging drops as before at ambient temperature (22°C) using the factorial crystallization screening trials of Jancarik & Kim (1991) and using the commercially available 30 product from Hampton Research, Riverside). The following are optimum conditions: hexagonal bipyrimidal crystals are grown at 4°C for 2-3 days from hanging drops containing 1.0-1.2 M sodium acetate (pH unadjusted) and 0.1 M sodium cacodylate (pH 7.4), 3 mM DTT, with either a 1 μl protein solution, 1 μl precipitant solution or 2

µl protein solution, 1 µl precipitant solution and a 0.6 ml reservoir using silanized coverslip, at a protein concentration of 7-10 mg/ml. The best crystals have a limiting dimension of 200 μm. The following are optimum conditions for crystallization of the TR-β LBD with GC-1: hexagonal bipyrimidal crystals are grown at 4°C for 2-3 days from hanging drops containing 0.8-1.0M sodium acetate (pH unadjusted), 50-200nM sodium succinate, and 0.1M sodium cacodylate (pH 7.2), 3mM DTT, 1 μl protein solution, 1 μl precipitant solution and a 0.6ml reservoir using silanized coverslip, at a protein concentration of 7-10 mg/ml. The best crystals have a limiting dimension of 200μM. The unit cell dimensions are cell length a=b=68.73, cell length 10 c=130.09. The unit cell angles are α=90°, β=90°, γ=120°.

The crystal system for human TR- $\beta$  LBD bound to T<sub>3</sub>, Triac or GC-1 is trigonal with the space group p3<sub>1</sub>21. The unit cell dimensions are cell length a = cell length b = 68.448 angstroms, cell length c = 130.559 angstroms. The angles are  $\alpha$  = 90°,  $\beta$  = 90°, gamma = 120°.

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# Example 9 - Determination of Liganded TR- $\alpha$ LBD and TR- $\beta$ Crystal Structures

Data from each cocrystal (Rat TR-α LBD with Dimit, T3 and IpBr2; Human TR-β LBD with Triac and GC-1) is measured on a Mar area detector at Stanford 20 Synchrotron Radiation Laboratory beamline 7-1 (λ = 1.08 angstroms) using 1.2° oscillations. Data from the cocrystal of the hTR-β LBD with Triac is measured on a Mar area detector at Stanford Synchrotron Radiations Laboratory beamline 7-1 (λ = 1.08 angstroms) using 1.0 oscillations. Data from the cocrystal of the hTR-β LBd with GC-1 is measured on a R-axis II area detector on a Rigaku rotating Cu anode 25 (50kV, 300mA). The crystals are transferred into a cryosolvent containing 1.2M sodium acetate, 0.1M sodium cacodylate, adn 15% glycerol followed by a second transfer into 30% glycerol, then flash frozen in liquid nitrogen. An orientation matrix for each crystal is obtained using DENZO. The reflections are integrated with DENZO (commercially available from Molecular Structure Corp., The Woodlands, 30 Texas) and are scaled with SCALEPACK (as described in Otwinowski, Z, *Proceedings of the CCP4 Study Weekend: "Data Collection and Processing,"* 56-62 (SERC Daresbury Laboratory, Warrington, UK 1993) incorporated by reference).

For rTR-α cocrystals, data from the T<sub>3</sub> cocrystal is measured with the b\* axis approximately parallel with the spindle. The crystals are flash frozen at -178°C in a nitrogen gas stream with the MPD mother liquor serving as the cryosolvent. An orientation matrix for each crystal is determined using REFIX (Kabsch, W., J. Appl. 5 Crystallogr. (1993) 26:795-800 incorporated by reference). Reflections are integrated with DENZO, and are scaled with SCALEPACK.

For the T<sub>3</sub> data set, Bijvoet pairs are kept separate, and are locally scaled using MADSYS (W. Hendrickson (Columbia University) and W. Weis (Stanford University)).

10 Cocrystals prepared from the three isosteric ligands are isomorphous. MIR analysis is performed using programs from the CCP4 suite (Collaborative Computational Project, N.R. Acta Crystallogr. (1994) D50:760-763, incorporated by reference herein). Difference Pattersons is calculated for both T<sub>3</sub> and IpBr<sub>2</sub>, taking the Dimit cocrystal as the parent. The positions of the three iodine atoms in the T<sub>3</sub> 15 difference Patterson are unambiguously determined from the Harker section of the density map as peaks of 11[ above background. The positions for the two bromine atoms in the IpBr2 cocrystals, are located independently, as peaks 8[ above the noise level. Phases for the LBD-122/410 are calculated from the solution to the IpBr<sub>2</sub> difference Patterson, and are used to confirm the location of the unique third iodine of 20 the T<sub>3</sub> cocrystal. Halogen positions are refined with MLPHARE, including the anomalous contributions from the iodine atoms (Otwinowski, Z, Proceedings of the CCPR Study Weekend 80-86 (SERC Daresbury Laboratory, Warrington, UK 1991)). The MIRAS phases are improved through solvent flattening/histogram matching using DM (Cowtan, K., Joint CCP4 and ESF-EACBM Newsletter on Protein 25 Crystallography (1994) 31: 34-38, incorporated by reference herein).

A model of the LBD-122/410 with Dimit bound is built with the program O from the solvent flattened MIRAS 2.5 angstrom electron density map (Jones et al., Acta Crystallogr. (1991) A 47:110-119, incorporated by reference herein). The initial model, without ligand, (Rcryst = 40.1%), is refined using least-squares protocols with XPLOR. The Dimit ligand is built into unambiguous Fo-Fc difference density during the following round. Subsequent refinement employs both least-squares and simulated annealing protocols with XPLOR (Brunger et al., Science (1987) 235:458-460), incorporated by reference herein). Individual atomic B-factors are refined

Page 81 of 447

isotropically. As defined in PROCHECK, all residues are in allowed main-chain torsion angle regions as described in Laskowski *et al.*, *J. Appl. Crystallogr.*, (1993) 26:283-291, incorporated by reference herein. The current model is missing 34 residues (Met<sub>122</sub>-Gln<sub>156</sub>) at the N-terminus, and 5 residues (Glu<sub>406</sub>-Val<sub>410</sub>) at the C-5 terminus.

In addition, the following residues are not modeled beyond Cβ due to poor density: 184, 186, 190, 198, 206, 209, 240, 301, 330, 337, 340, 343, 359, and 395. The average B-value for protein atoms is 34.5 Å<sup>2</sup>. The final model consists of the LBD-122/410, residues Arg<sub>157</sub>-Ser<sub>183</sub>, Trp<sub>185</sub>-Gly<sub>197</sub>, Ser<sub>199</sub>-Asp<sub>206</sub> and Asp<sub>208</sub>-Phe<sub>405</sub>; 10 three cacodylate-modified cysteines: Cys<sub>334</sub>, Cys<sub>380</sub> and Cys<sub>392</sub>; and 73 solvent molecules modeled as water (2003 atoms).

\*
$$R_{\text{sym}} = 100 \times \Sigma_{\text{hkl}} \Sigma_{\text{i}} \mid I_{\text{i}} - I \mid / \Sigma_{\text{hkl}} \Sigma_{\text{i}} I_{\text{i}}$$
  
† $R_{\text{der}} = 100 \times \Sigma_{\text{hkl}} \mid F_{\text{PH}} - F_{\text{H}} \mid / \Sigma_{\text{hkl}} \mid F_{\text{P}} \mid$ 

The occupancy for the two bromine sites is set to 35 electrons. The occupancies of the 15 iodine sites are relative to this value.

§Phasing power =  $\langle FH \rangle$ ,  $/ \langle \epsilon \rangle$ , where  $\langle FH \rangle$  is the mean calculated heavy atom structure factor amplitude and  $\langle \epsilon \rangle$  is the mean estimated lack of closure.

4Rcullis =  $\langle \in \rangle / \langle iso \rangle$ , where  $\langle \in \rangle$  is the mean estimated lack of closure and  $\langle iso \rangle$  is the isomorphous difference.

20 ¶Rcryst = 100 x  $\Sigma_{hkl}$  |F<sub>o</sub>-Fc| /  $\Sigma_{hkl}$  |F<sub>o</sub>| where F<sub>o</sub> and F<sub>c</sub> are the observed and calculated structure factor amplitudes (for data F/ $\sigma$  > 2). The Rfree was calculated using 3% of the data, chosen randomly, and omitted from the refinement.

§ Correlation coefficient = 
$$\Sigma_{hkl}$$
 ( $|F_o| - |F_o|$ ) x ( $|F_c| - |F_c|$ )/ $\Sigma_{hkl}$  ( $|F_o| - |F_o|$ )<sup>2</sup>x $\Sigma_{hkl}$  ( $|F_c| - |F_c|$ )<sup>2</sup>

25

# Example 10. Phasing of the rTR- $\alpha$ LBD and hTR- $\beta$ LBD complex with Triac

Due to the possible non-isomorphism of the rTRα LBD complex with Triac, a molecular replacement solution is determined using AMORE (Navaza, J., Acta 30 Crystallographica Section A-Fundamentals of Crystallography (1994) 50:157-63 from a starting model consisting of rTRI LBD complex with T<sub>3</sub>, but with the ligand, all water molecules, and the following residues omitted: Asn 179, Arg228, Arg262, Arg266, and Ser 277. Strong peaks are obtained in both the rotation and translation

searches, with no significant (> 0.5 times the top peak) false solutions observed (Table 6). Strong positive density present in both the anomalous and conventional difference Fourier maps confirm the solution. Maps are calculated using sigma-A weighted coefficients output by REFMAC (Murshudov, et al. "Application of 5 Maximum Likelihood Refinements," in Refinement of Protein Structures, Proceedings of Daresbury Study Weekend (1996)) after 15 cycles of maximum likelihood refinement. Triac, the omitted residues, and water molecules 503, 504, 534 (following the numbering convention for the TR complex with T3) are built into the resulting difference density using O (Jones et. al.); the conformations of these residues are further confirmed in a simulated-annealing omit map (Brunger et. al.). The complete model is then refined using positional least-squares, simulated annealing, and restrained, grouped B factor refinement in XPLOR to an Reryst of 23.6% and an Rfree of 24.1%

Phasing of a related LBD using the structure of the rTR-α LBD is conducted 15 as follows. A molecular replacement solution for the hTR-β LBD complex with Triac is determined using AMORE from a starting model consisting of the rTR-a LBD complexed with T3, but with the ligand and all water molecules omitted. Strong peaks are obtained in both the rotation and translation searches, with no significant (>0.5 times the top peak) false solutions (Table 7). Strong positive density present in 20 both the anomalous and conventional difference Fourier maps confirm the solution. Initial maps are calulated using sigma-A weighted coefficients output by REFMAC after 9 cycles of maximum likelihood refinement. The real-space fit for each residues was calculated using OOPS (Kleywegt, GJ and Jones, TA, OOPS-a-daisy, ESF/CCP4 Newsletter 30, June 1994, pp. 20-24) and the residues with a real-space fit less than 2 25 standard deviations below the mean removed: Ala253-Lys263; Glu245-Leu250. To reduce bias, the following residues were modeled as alanine: Arg282, Arg316, Arg 320, Asn 331: Cycles of rebuilding and positional least-squares, simulated annealing, and restrained, grouped B factor refinement with XPLOR produce a model with an R<sub>cryst</sub> of 25.3 and an R<sub>free</sub> of 28.9%. The final model consists of hTR-β LBD residues 30 Glu202-Gln252, Val264-Glu460; three cacodylate-modified cysteines with the cacodylate moeity modeled as free arsenic: Cys294, Cys298, Cys388, and Cys434; and 35 solvent molecules modeled as water.

# EXAMPLE 11. CONNECTING QSAR WITH STRUCTURE IN THE THYROID HORMONE RECEPTOR

The conclusions of classic thyroid hormone receptor quantitative structureactivity relationships may be summarized as follows:

- 5 1) the R<sub>4</sub>'-hydroxyl group functions as a hydrogen bond donor;
  - 2) the amino-propionic acid interacts electrostatically through the carboxylate anion with a positively charged residue from the receptor;
    - 3) the preferences of  $R_3/R_5$  substituent are I>Br>Me>>H;
    - 4) the preferences of the R<sub>3</sub>'-substituent are Ipr>I>Br>Me>>H.
- 10 The structure of the thyroid hormone receptor ligand binding domain complexed with the agonists T3, IpBr<sub>2</sub>, Dimit, Triac, and GC1 as provided herein, permits:
  - the identification of receptor determinants of binding at the level of the hydrogen bond;
  - the association of these determinants with the predictions of classic thyroid hormone receptor QSAR; and
    - 3) prediction as to which determinants of binding are rigid, and which are flexible, for both the ligand and the receptor.

This classification for the agonists of the type ( $R_1$ =amino-propionic, acetic acid;  $R_3$ , $R_5$ =I, $B_7$ ,Me;  $R_3$ '=Ipr,I) is given below (for the representative ligand  $T_3$ );

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15

F = Fiducial (always satisfied)

A = Adjustable

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Based upon the methods and data described herein, the following is an embodiment of the computational methods of the invention, which permit design of nuclear receptor ligands based upon interactions between the structure of the amino acid residues of the receptor LBD and the four different ligands described herein. The

small molecule structures for the ligands can be obtained from Cambridge Structural Database (CSD), and three dimensional models can be constructed using the methods described throughout the specification. The following are factors to consider in designing synthetic ligands:

- 1) Histidine 381 acts as a hydrogen bond acceptor for the R<sub>4</sub>' hydroxyl, with the optimal tautomer maintained by water molecules. See FIG. 23 and FIG 24. Histidine is the only hydrophilic residue in this hydrophobic pocket that surrounds the R<sub>4</sub>' substituent. Histidine can be either a hydrogen bond acceptor or donor, depending on its tautomeric state. It is preferably a hydrogen bond donor, but can tolerate being a hydrogen bond acceptor, as for example, when there is a methoxy at the R<sub>4</sub>' position of the ligand;
- Arginines 228, 262, and 266 interact directly and through water-mediated hydrogen bonds with the R<sub>1</sub>-substituent, with the electrostatic interaction provided by Arginine 266 (as in the Triac complex). This polar pocket is illustrated by FIG. 23 -15 FIG. 25. FIG. 23 depicts T<sub>3</sub> in the TRI ligand binding cavity, where T3's aminopropionic R<sub>1</sub>- substituent interacts with Arg 228, HOH502, H9H503 and HOH504 via hydrogen bonds. FIG. 24 depicts Triac in the ligand binding cavity, with its -COOH R<sub>1</sub> substituent in the polar pocket. In FIG. 24, Arg 228 no longer shares a hydrogen bond with the ligand, but the -COOH R1 substituent forms hydrogen bonds with Arg 20 266. FIG. 25 superimposes T<sub>3</sub> and Triac in the ligand binding cavity and shows several positionally unchanged amino acids and water molecules, and selected changed interacting amino acids and water molecules. The three figures illustrate parts of the polar pocket that can change and those parts that do not move upon binding of different ligands. For example, the Arg 262 at the top of the polar pocket 25 does not move, even when the R<sub>1</sub> substituent has changed from a -COOH to an aminopropionic acid group. However, the other two Arginines, Arg 228 and Arg 266, demonstrate flexibility in the polar pocket to respond to the change in the size or chemical naure of the R<sub>1</sub> substituent.
- 3) Inner and outer pockets for the R<sub>3</sub>/R<sub>5</sub> substituents are formed by 30 Ser260, Ala263, Ile299; and Phe 218, Ile221, Ile222, respectively. See FIGS. 21 and 22. The inner pocket is filled by either the R<sub>3</sub> or the R<sub>5</sub> substituent, regardless of the size of the substituent, and may act as a binding determinant by positioning the ligand in the receptor. Optimally, the inner pocket amino acids interact with an R<sub>3</sub> or R<sub>5</sub>

substituent that is no larger than an iodo group. If the inner pocket is filled by the R<sub>3</sub> substituent, then the outer pocket interacts with the R<sub>5</sub> substituent and vice versa. The outer pocket can adjust to the size of its substituent through main chain motion centered at the break in helix 3 (Lys220-Ile221), suggesting that the bending of H3, and motion of the N-terminal portion of H3, may represent a conformational change induced on ligand binding. The outer pocket has greater flexibility than does the inner pocket in terms of accommodating a larger substituent group.

4) A pocket for the R<sub>3</sub>'-substituent is formed by Phe 215, Gly290, Met388. The pocket is incompletely filled by the R<sub>3</sub>'-iodo substituent, and 10 accommodates the slightly larger 3'-isopropyl substituent by movement of the flexible Met388 side chain and the H7/H8 loop. This pocket can accommodate R<sub>3</sub>' substituents that are even larger than isopropyl, for example, a phenyl group.

The above information will facilitate the design of high affinity agonists and antagonists by improving automated QSAR methodologies and informing manual modeling of pharmaceutical lead compounds. For example, the inclusion of discrete water molecules provides a complete description of hydrogen bonding in the polar pocket for use with pharmacophore development: also, the identification of mobile and immobile residues within the receptor suggests physically reasonable constraints for use in molecular mechanics/dynamics calculations.

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#### EXAMPLE 12. DESIGN OF AN INCREASED AFFINITY LIGAND

Direct interaction between the receptor and the ligand is limited in the polar pocket, which interacts with the R<sub>1</sub> substituent. While the lack of complementarity may contain implications for biological regulation, it also provides an opportunity for increasing affinity by optimizing the interaction between the amino acids of the polar pocket and the R<sub>1</sub> substituent of a synthetic ligand. The structure of the receptor-ligand interactions described herein enables design of an increased affinity synthetic ligand having two complementary modifications:

1) Remove the positively charged amine. The strongly positive 30 electrostatic potential predicted for the polar pocket suggests that the positively charged amine of the aminopropionic acid R<sub>1</sub> substituent may be detrimental to binding. Suitable groups for substitution are suggested by the nature of nearby hydrogen bond partners: for example, Thr 275 O or Ser 277 N. See e.g. Tables in Appendix 2. For example, any any negatively charged substituent would be

Page 86 of 447

WO 99/26966 PCT/US98/25296

compatible for interacting with the amino acids of the polar pocket, including carboxylates, carbonyl, phosphonates, and sulfates, comprising 0 to 4 carbons. Another example of an R<sub>1</sub> substitution is an oxamic acid that replaces the amine of the naturally occurring ligand with one or more carbonyl groups.

5 Incorporate hydrogen bond acceptor and donor groups into the R<sub>1</sub>-2) substituent to provide broader interactions with the polar pocket scaffold. Such hydrogen bond acceptor and donor groups incorporated into the R1-substituent will allow interactions that would otherwise occur with water molecules in the polar pocket. Specific waters include HOH 504 (hydrogen bonds with Ala 225 O and Arg 10 262 NH); and HOH 503 hydrogen bonds with Asn 179 OD1, Ala 180 N), both of which are present in all four complexes (TR LBD complexed with T3, TR LBD complexed with IpBr2, TR LBD complexed with Dimit and TR LBD complexed with Triac). Analysis of the hydrogen bonding network in the polar pocket suggests replacement of HOH 504 with a hydrogen bond acceptor, and HOH 503 with an 15 hydrogen bond donor (although the chemical nature of asparagine probably permits flexibility at this site). Thus, incorporating a hydrogen bond acceptor in an R1 substituent that could take the place of the HOH504 or incorporating a hydrogen bond acceptor in an R1 substituent that could positionally replace the HOH503, or a combination thereof, are methods of designing novel synthetic TR ligands.

These two design approaches can be used separately or in combination to design synthetic ligands, including those in Table 5 (below).

A corollary to this approach is to design specific interactions to the residues Arg262 and Asn 179. The goal is to build in interactions to these residues by designing ligands that have R<sub>1</sub> substituents that form hydrogen bonds with water 25 molecules or charged residues in the polar pocket.

High-affinity ligands also may be designed and selected using small molecules that bind to proximal subsites of the target nuclear hormone receptor that are identified in a structure-based screen and then linked together in their experimentally determined bound orientiations. Such a method has been described in design of high-affinity ligands for the FK506 binding protein (FKBP), stromelysin, gelatinase A, and human papillomavirus E2 (Hajduk et al., Science 278:497-499 (1997)), which reference and its references are incorporated herein by reference. The preferred small molecules for screening are compounds of Formula I or derivatives thereof. For

example, a compound of Formula I (φ-X-φ) or a derivative thereof (φ-X or X-φ) is screened for binding a target nuclear hormone receptor LBD. Proximal subsites of the nuclear hormone receptor include the hydrophobic and polar pockets of the LBD, and substites extended therefrom. As an example, Fourier transformation or nuclear 5 magnetic resonance (NMR) -based structure screens can be used. When a NMR-based screen is used, binding can be detected from the amide chemical shift changes observed in two-dimensional heteronuclear single quantum correlation (HSQC) spectra aquired in the presence and absence of added compound. Once two ligands are identified that bind to the receptor, the crystal or solution structure of the ternary 10 complex is determined. From the structural information, a compound is synthesized which links the two ligands, where the linker is selected based on structural information. The new compound is then screened for binding affinity, for example, using a binding assay as described herein. Only a few linked ligands need to synthesized and screened when using this approach.

15 Compounds of the invention also may be interatively designed from structural information of the compounds described above using other structure-based design/modeling techniques (Jackson, R.C., Contributions of protein structure-based drug design to cancer chemotherapy. Semninars in Oncology, 1997, 24(2)L164-172; and Jones, T.R., et al., J. Med. Chem., 1996 39(4):904-917).

WO 99/26966

PCT/US98/25296

Table 5: Synthetic TR Ligands

5

R1	R2	R3	R5	R6	x	R'2	R'3	R'4	R'5	R'6
СО2Н	Н	Me <sub>.</sub>	Me	Н	0	Н	Me	ОН	Me	Н
СН2СО2Н		1	1		s		Et	SH	Et	
СН2СН2СО2Н		Br	Br				nPr	NH2	nPr	
CH2CH(NH2)CO2H		Cl	CI				iPr		iPr	
ОСН2СО2Н		Et	Et				Ph		nBu	
ОСН2СН2СО2Н		OH-	ОН				I		nPen	
NHCH2CO2H		NH2	NH2				Br	·	nHex	
NHCH2CH2CO2H		SH	SH				Cl		Ph	
СН2СОСОСО2Н									hetero	
•									cycle	
NHCOCOCO2H									aryl	
СОСО2Н										
CF2CO2H										
COCH2CO2H										

Any combination of the above substituents in the biphenyl ether scaffold structure shown above may result in a potentially pharmacologically useful ligand for the 10 thyroid hormone receptor. These novel ligands may be antagonists of the thyroid receptor.

WO 99/26966

PCT/US98/25296

**TABLE 6: TR-α LBD-122/410** 

				<b></b>
	Dimit '	Т3	lpBr₂	Triac
Data collection				
Cell dimensions				
a (Å)	117.16	117.19	117.18	118.19
b (Å)	80.52	80.20	80.12	81.37
c (Å)	63.21	63.23	63.13	63.73
β (°)	120.58	120.60	120.69	121.00
Resolution (Å)	2.2	2.0	2.1	2.45
Obs. Reflections, (no.)	57031	64424	66877	83573
Unique Reflections, (no.)	22327	21023	23966	18453
Completeness, (%)	87.0	82.4	93.7	96.0
*R <sub>svm</sub> (%)	3.9	3.5	4.5	7.5
Phasing (15.0 - 2.5Å)				
†R <sub>der</sub> (%)	· -	19.6	11.6	
No. of sites	-	3	2	
‡Occupancy	-	44.6 (19.8)	35.0	
(Anomalous)	-	50.2 (23.7)	35.0	
		39.2 (22.3)		
§F <sub>H</sub> /E				
centric (acentric)		•		
15.0-5.0 Å	•	3.67 (4.61)	2.25 (3.09)	
5.0-3.0 Å	-	2.23 (2.75)	1.25 (1.85)	
3.0-2.5 Å	•	1.64 (1.99)	1.15 (1.57)	
¶R <sub>Cullis</sub> (%)				
15.0-5.0 Å	-	33	44	
5.0-3.0 Å	-	45	63	
3.0-2.5 Å	•	. 60	65	
Mean figure of merit	0.62	-	•	
MR Phasing (10-3.5Å)				
Rotation Search:	•			$\Theta_1 = 309.37$
Euler Angles (°)				$\Theta_2 = 48.96$
				$\Theta_3 = 127.28$
§ correlation coefficient				34.3
Translation Search: Fractional coordinates				x = 0.1571
				y = 0.000
				z = 0.3421
§ correlation Coefficient				65.8
<sup>1</sup> R factor		•		31.2

Refinement Resolution (Å)	15.0-2.2	5.0 - 2.0	15.0 - 2.2	25-2,5
¶R <sub>cryst (%)</sub>	20.5	22.1	21.4	23.6
R <sub>free (%)</sub>	22.7	24.0	22.4	24.1

**TABLE 7: TR-β LBD-202/461** 

	Triac	Т3	GCI
Data collection			
Space Group	<del></del>	P3121	
Cell dimensions			
a (Å)	68.9	68.45	68.73
c (Å)	131.5	130.56	130.09
Resolution (Å)	2.4	3.1	2.8
Obs. Reflections. (no.)	80196	55103	54104
Unique Reflections. (no.)	14277	6847	8987
Coverage (%)	97.0	95.7	97.1
*R <sub>sym</sub> (%)	5.1	4.6	5.5
MR Phasing (15.0 - 3.5Å)			
Rotation Search	$\Theta_1 = 39.13$		
Euler Angles (°)	Θ <sub>2</sub> =68:00		
	⊕₃=323.6		
§ correlation coefficient	21.6		
(Highest false peak)	(10.8)		
Translation Search	x=0.748		
Fractional Coordinates	y=0.158		
	z=0.167		
§ correlation coefficient	57.5		
(Highest false peak)	(38.7)		
	0.612		
*R factor	40.7	40.8	
Refinement			
Resolution (Å)	30-2.4		30-2.9
¶R <sub>cryst (%)</sub>	25.3		27.3
R <sub>free (%)</sub>	28.9		33.4

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference. The nuclear receptor ligands, particularly the TR ligands, of these

references are herein incorporated by reference and can be optionally excluded from the claimed compounds with a proviso.

Headings and subheadings are presented only for the convenience of the reader and should not be used to construe the meaning of terms used within such 5 headings and subheadings.

The invention now being fully described, it will be apparent to one of ordinary skill in the art that many changes and modifications can be made thereto without departing from the spirit or scope of the appended claims.

#### WHAT IS CLAIMED IS:

A method of modulating the activity of a thyroid hormone receptor
 (TR) which comprises administering to a mammal in need thereof a compound of the
 formula:

wherein said compound fits spatially and preferentially into a TR ligand binding domain (TR LBD) and comprises the following substituents:

- (i) an R<sub>1</sub>-substituent comprising an anionic group that interacts with a side 15 chain nitrogen atom of an arginine corresponding to a residue selected from the group consisting of Arg228, Arg262, and Arg266 of human TR-α, and Arg282, Arg316 and Arg320 of human TR-β, and wherein the anionic group is 1.7-4.0Å from the nitrogen atom;
- (ii) an R<sub>2</sub>-substituent comprising a hydrophobic or hydrophilic group that
   20 fits spacially into the TR LBD;
- (iii) an R<sub>3</sub>-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a serine, alanine or isoleucine corresponding to a residue selected from the group consisting of Ser260, Ala263 and Ile299 of human TR-α, and Ser314, Ala317 and Ile352 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;

(iv) an R<sub>5</sub>-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a phenylalanine or isoleucine corresponding to a residue selected from the group consisiting of Phe218, Ile221 and Ile222 of human TR-I, and Phe272, Ile275 and Ile276 of human TR-β, and wherein the hydrophobic or bydrophilic group is 1.7-4.0Å from the side chain atom;

- (v) an R<sub>6</sub>-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- (vi) an X-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a leucine corresponding to a residue selected from the group consisting of Leu276 and Leu292 of human TR-α, and Leu 330 and Leu346 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;
  - (vii) an R<sub>2</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- 15 (viii) an R<sub>3</sub>'-substituent comprising a hydrophobic group that interacts with a side chain atom of a phenylalamine, glycine or methionine corresponding to a residue selected from the group consisting of Phe215, Gly290, and Met388 of human TR-α, and Phe269, Gly344, Met442 of human TR-β, and wherein the hydrophobic group is 1.7-4.0Å from the side chain atom;
- 20 (ix) an R<sub>4</sub>'-substituent comprising an hydrogen bond donor or acceptor group that interacts with a side chain carbon or nitrogen atom of a histadine corresponding to residue His381 of human TR-α, and His435 of human TR-β, and wherein the hydrogen bond donor or acceptor group is 1.7-4.0Å from the side chain atom;

(x) an R<sub>5</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;

- (xi) and R<sub>6</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- wherein said compound is other than a thyronine or thyronine-like compound disclosed in a reference cited in Appendix I, and wherein the activity of said TR is modulated.
  - 2. The method according to claim 1,
- 10 wherein R<sub>1</sub> is
  - -O-CH2CO2H, -NHCH2CO2H,
  - -CO<sub>2</sub>H, -CH<sub>2</sub>CO<sub>2</sub>H, -CH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>H, -CH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>H,
  - -CH<sub>2</sub>CH(NH<sub>2</sub>)CO<sub>2</sub>H,

-CH<sub>2</sub>CH[NHCOCH $\phi_2$ ]CO<sub>2</sub>H,

- -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>
- ]CO₂H,
- -CH<sub>2</sub>CH[NH-FMOC]CO<sub>2</sub>H,
- -CH<sub>2</sub>CH[NH-tBOC]CO<sub>2</sub>H, or a carboxylate connected to the ring with a 0 to 3 carbon linker,
  - $-PO_{3}H_{2}, \qquad -CH_{2}PO_{3}H_{2}, \qquad -CH_{2}CH_{2}PO_{3}H_{2}, \qquad -CH_{2}CHNH_{2}PO_{3}H_{2},$
  - $-CH_2CH[NHCOCH_{\phi_2}]PO_3H_2, \qquad -CH_2CH[NHCO(CH_2)_{15}CH_3]PO_3H_2, \\$
- -CH<sub>2</sub>CH[NH-FMOC]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub> CH[NH-tBOC]PO<sub>3</sub>H<sub>2</sub>, or a phosphate or phosphonate connected to the ring with a 0 to 3 carbon linker,

-CH<sub>2</sub>CH[NH-FMOC]SO<sub>3</sub>H, -CH<sub>2</sub> CH[NH-tBOC]SO<sub>3</sub>H, or a sulfate or sulfite connected to the ring with a 0 to 3 carbon linker,

or acts as the functional equivalent of CH<sub>2</sub>CH(NH<sub>2</sub>)CO<sub>2</sub>H of T3 in the molecular recognition domain when bound to a TR, wherein said R<sub>1</sub> can be optionally substituted with an amine,

#### wherein R2 is

H, halogen, CF<sub>3</sub>, OH, NH<sub>2</sub>, SH, CH<sub>3</sub>, -Et,

or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

#### wherein R<sub>3</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et,

or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

#### wherein R5 is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR, and R<sub>3</sub> can be identical to R<sub>5</sub>,

## wherein R<sub>6</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR, and R<sub>2</sub> can be identical to R<sub>6</sub>,

## 5 wherein R<sub>2</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

wherein R<sub>3</sub>' is any hydrophobic group, including

halogen, -CF<sub>3</sub>, -SH, alkyl, aryl, 5- or 6-membered heterocyclie, cyano, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

#### wherein R<sub>4</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, -SH, -CH<sub>3</sub>, -Et, or akyl, aryl or 5- or 6-membered heterocyclic aromatic attached through urea or carbamate linkages to O or N or S at the R<sub>4</sub>' position, or acts as the functional equivalent of OH in the molecular recognition domain when bound to a TR,

20

25

## wherein R5' is

-H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally

connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups,

wherein R<sub>6</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

wherein X is

O, S, SO<sub>2</sub>, NH, NR<sub>7</sub>, CH<sub>2</sub>, CHR<sub>7</sub>, CR<sub>7</sub>R<sub>7</sub>, wherein R<sub>7</sub> is alkyl, aryl or 5- or 6-membered heterocyclic aromatic,

15

5

and wherein said TR LBD ligand has an apparent Kd for binding TR LBD of 1 TM or less.

- 3. The method of claim 2, wherein
- $R_1$  is carboxylate, phosphonate, phosphate or sulfite and is connected to the ring with a 0 to 3 carbon linker,

R<sub>2</sub> is H,

 $R_3$  is -I, -Br, or -CH<sub>3</sub>,

 $R_5$  is -I, -Br, or -CH<sub>3</sub>,

25 R<sub>6</sub> is H,

R<sub>2</sub>' is H,

R<sub>3</sub>' is -I, -Br, -CH<sub>3</sub>, -iPr, -phenyl, benzyl, or 5- or 6-membered ring heterocycles,

R<sub>4</sub>' is -OH, -NH<sub>2</sub>, and -SH,

R<sub>5</sub>' is -H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups, and

 $R_6$  is H.

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- 4. The method of claim 1, wherein said compound fits spatially and preferentially into TR LBD isoform  $\alpha$  (TR- $\alpha$ ).
- 5. The method of claim 4, wherein said compound comprises an anionic group that interacts with the side chain oxygen or carbon of a serine residue corresponding to Ser277 of human TR-α, and wherein the anionic group is 1.7-4.0Å from the side chain atom.

6. The method of claim 1, wherein said compound fits spatially and preferentially into TR LBD isoform  $\beta$  (TR- $\beta$ ).

- The method of claim 6, wherein said compound comprises an anionic
   group that interacts with the side chain nitrogen of an arginine corresponding to
   Asn331 of human TR-β, and the anionic group is 1.7-4.0Å from the side chain atom.
- 8. A method for identifying a compound capable of selectively modulating the activity of a thyroid hormone receptor (TR) isoform, said method 10 comprising:

modeling test compounds that fit spacially and preferentially into a TR ligand binding domain (TR LBD) isoform of interest using an atomic structural model of a TR LBD isoform bound to a test compound,

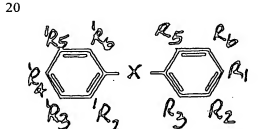
screening said test compounds in a biological assay for TR isoform

15 activity characterized by binding of a test compound to a TR LBD isoform, and

identifying a test compound that selectively modulates the activity of a

TR isoform.

9. The method of claim 8, wherein said compound is of the formula:



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which comprises the following substituents:

- (i) an R<sub>1</sub>-substituent comprising an anionic group that interacts with a side chain nitrogen atom of an arginine corresponding to a residue selected from the group consisting of Arg228, Arg262, and Arg266 of human TR-α, and Arg282, Arg316 and Arg320 of human TR-β, and wherein the anionic group is 1.7-4.0Å from the nitrogen atom;
  - (ii) an R<sub>2</sub>-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- (iii) an R<sub>3</sub>-substituent comprising a hydrophobic or hydrophilic group that 10 interacts with a side chain atom of a serine, alanine or isoleucine corresponding to a residue selected from the group consisting of Ser260, Ala263 and Ile299 of human TR-α, and Ser314, Ala317 and Ile352 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;
- (iv) an R<sub>5</sub>-substituent comprising a hydrophobic or hydrophilic group that
  15 interacts with a side chain atom of a phenylalanine or isoleucine corresponding to a residue selected from the group consisiting of Phe218, Ile221 and Ile222 of human TR-α, and Phe272, Ile275 and Ile276 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;
- (v) an R<sub>6</sub>-substituent comprising a hydrophobic or hydrophilic group that
   20 fits spacially into the TR LBD;
- (vi) an X-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a leucine corresponding to a residue selected from the group consisting of Leu276 and Leu292 of human TR-α, and Leu 330 and Leu346 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from 25 the side chain atom;

(vii) an R<sub>2</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;

- (viii) an R<sub>3</sub>'-substituent comprising a hydrophobic group that interacts with a side chain atom of a phenylalamine, glycine or methionine corresponding to a residue selected from the group consisting of Phe215, Gly290, and Met388 of human TR-α, and Phe269, Gly344, Met442 of human TR-β, and wherein the hydrophobic group is 1.7-4.0Å from the side chain atom;
- (ix) an R<sub>4</sub>'-substituent comprising an hydrogen bond donor or acceptor group that interacts with a side chain carbon or nitrogen atom of a histadine 10 corresponding to residue His381 of human TR-α, and His435 of human TR-β, and wherein the hydrogen bond donor or acceptor group is 1.7-4.0Å from the side chain atom;
  - (x) an R<sub>5</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD; and
- 15 (xi) and R<sub>6</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD.
  - 10. The method according to claim 9, wherein  $R_1$  is
- 20 -O-CH<sub>2</sub>CO<sub>2</sub>H, -NHCH<sub>2</sub>CO<sub>2</sub>H,
  - -CO<sub>2</sub>H, -CH<sub>2</sub>CO<sub>2</sub>H, -CH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>H, -CH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>H,
  - -CH<sub>2</sub>CH(NH<sub>2</sub>)CO<sub>2</sub>H,

- $CH_2CH[NHCOCH\phi_2]CO_2H$ ,

- -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>
- ]CO<sub>2</sub>H,
- -CH2CH[NH-FMOC]CO2H,
- -CH<sub>2</sub>CH[NH-tBOC]CO<sub>2</sub>H, or a carboxylate connected to the ring with a 0 to 3
- 25 carbon linker,

WO 99/26966

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PCT/US98/25296

-PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CHNH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>,
-CH<sub>2</sub>CH[NHCOCHφ<sub>2</sub>]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>]PO<sub>3</sub>H<sub>2</sub>,
-CH<sub>2</sub>CH[NH-FMOC]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub> CH[NH-tBOC]PO<sub>3</sub>H<sub>2</sub>, or a phosphate or phosphonate connected to the ring with a 0 to 3 carbon linker,

-SO<sub>3</sub>H, -CH<sub>2</sub>SO<sub>3</sub>H, -CH<sub>2</sub>CH<sub>2</sub>SO<sub>3</sub>H, -CH<sub>2</sub>CHNH<sub>2</sub>SO<sub>3</sub>H,
-CH<sub>2</sub>CH[NHCOCHφ<sub>2</sub>]SO<sub>3</sub>H, -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>]SO<sub>3</sub>H,
-CH<sub>2</sub>CH[NH-FMOC]SO<sub>3</sub>H, -CH<sub>2</sub> CH[NH-tBOC]SO<sub>3</sub>H, or a sulfate or sulfite

10 connected to the ring with a 0 to 3 carbon linker,

or acts as the functional equivalent of CH<sub>2</sub>CH(NH<sub>2</sub>)CO<sub>2</sub>H of T3 in the molecular recognition domain when bound to a TR, wherein said R<sub>1</sub> can be optionally substituted with an amine,

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wherein R<sub>2</sub> is

H, halogen, CF<sub>3</sub>, OH, NH<sub>2</sub>, SH, CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

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wherein R<sub>3</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

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wherein R<sub>5</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR, and  $R_3$  can be identical to  $R_5$ ,

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wherein R<sub>6</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR, and R<sub>2</sub> can be identical to R<sub>6</sub>,

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wherein R2' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

15 wherein R<sub>3</sub>' is any hydrophobic group, including

halogen, -CF<sub>3</sub>, -SH, alkyl, aryl, 5- or 6-membered heterocyclie, cyano, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

### 20 wherein R<sub>4</sub>' is

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-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, -SH, -CH<sub>3</sub>, -Et, or akyl, aryl or 5- or 6-membered heterocyclic aromatic attached through urea or carbamate linkages to O or N or S at the R<sub>4</sub>' position, or acts as the functional equivalent of OH in the molecular recognition domain when bound to a TR,

wherein R5' is

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-H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups,

wherein R<sub>6</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

wherein X is

O, S, SO<sub>2</sub>, NH, NR<sub>7</sub>, CH<sub>2</sub>, CHR<sub>7</sub>, CR<sub>7</sub>R<sub>7</sub>, wherein R<sub>7</sub> is alkyl, aryl or 5- or 6-membered heterocyclic aromatic,

and wherein said TR LBD ligand has an apparent Kd for binding TR LBD of 1 TM or less.

25 11. The method of claim 10, wherein

 $R_1$  is carboxylate, phosphonate, phosphate or sulfite and is connected to the ring with a 0 to 3 carbon linker,

R<sub>2</sub> is H,

R<sub>3</sub> is -I, -Br, or -CH<sub>3</sub>,

 $R_5$  is -I, -Br, or -CH<sub>3</sub>,

5

R<sub>6</sub> is H,

R<sub>2</sub>' is H,

R<sub>3</sub>' is -I, -Br, -CH<sub>3</sub>, -iPr, -phenyl, benzyl, or 5- or 6-membered ring heterocycles,

10  $R_4$ ' is -OH, -NH<sub>2</sub>, and -SH,

R<sub>5</sub>' is -H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups, and

. R<sub>6</sub>' is H.

12. The method of claim 8, wherein said compound fits spatially and preferentially into TR LBD isoform  $\alpha$  (TR- $\alpha$ ).

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13. The method of claim 12, wherein said compound comprises an anionic group that interacts with the side chain oxygen or carbon of a serine residue corresponding to Ser277 of human TR- $\alpha$ , and wherein the anionic group is 1.7-4.0Å from the side chain atom.

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- 14. The method of claim 8, wherein said compound fits spatially and preferentially into TR LBD isoform  $\beta$  (TR- $\beta$ ).
- 15. The method of claim 14, wherein said compound comprises an anionic 10 group that interacts with the side chain nitrogen of an arginine corresponding to Asn331 of human TR-β, and the anionic group is 1.7-4.0Å from the side chain atom.
  - 16. The method of claim 8, wherein said compound binds to a TR LBD isoform with greater affinity than thyronine or triidothyronine.

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- 17. A method for identifying a thyroid hormone receptor (TR) agonist or antagonist ligand, said method comprising the steps of:
- providing the atomic coordinates of a TR ligand binding domain (TR LBD) to a computerized modeling system;
- 20 modeling ligands which fit spacially into the TR LBD; and
  - identifying in a biological assay for TR activity a ligand which increases or descreases the activity of said TR, whereby a TR agonist or antagonist is identified.

18. A peptide, peptidomimetic or synthetic molecule identified by the method of any one of claims 8 or 17, with the proviso that said molecule is other than a thyronine or thyronine-like compound disclosed in a reference cited in Appendix I.

5 19. A method of identifying a compound that selectively modulates the activity of a thyroid hormone receptor (TR) compared to other nuclear hormone receptors, said method comprising:

modeling compounds which fit spacially into a TR ligand binding domain (TR LBD) using an atomic structural model of a TR LBD,

selecting a compound comprising conformationally constrained structural features that interact with conformationally constrained residues of a TR LBD,

identifying in a biological assay for TR activity a compound that selectively binds to a TR LBD compared to other nuclear receptors, whereby a 15 compound that selectively modulates a TR is identified.

- 20. The method of claim 19, wherein said conformationally constrained residues of a TR LBD correspond to residues Met259, Leu276, Leu292, His381, Gly290, Ile221, and Phe401 of human TR-α, and residues Met313, Leu330, Leu346, 20 His435, Gly344, Ile275 and Phe455 of human TR-β.
  - 21. The method of claim 19, wherein said compounds are of the formula:

which comprises the following substituents:

- (i) an R<sub>1</sub>-substituent comprising an anionic group that interacts with a side chain nitrogen atom of an arginine corresponding to a residue selected from the group 10 consisting of Arg228, Arg262, and Arg266 of human TR-α, and Arg282, Arg316 and Arg320 of human TR-β, and wherein the anionic group is 1.7-4.0Å from the nitrogen atom;
  - (ii) an R<sub>2</sub>-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- 15 (iii) an R<sub>3</sub>-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a serine, alanine or isoleucine corresponding to a residue selected from the group consisting of Ser260, Ala263 and Ile299 of human TR-α, and Ser314, Ala317 and Ile352 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;
- 20 (iv) an R<sub>5</sub>-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a phenylalanine or isoleucine corresponding to a residue selected from the group consisting of Phe218, Ile221 and Ile222 of human TR-α, and Phe272, Ile275 and Ile276 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;

(v) an R<sub>6</sub>-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;

- (vi) an X-substituent comprising a hydrophobic or hydrophilic group that interacts with a side chain atom of a leucine corresponding to a residue selected from
  5 the group consisting of Leu276 and Leu292 of human TR-α, and Leu 330 and Leu346 of human TR-β, and wherein the hydrophobic or hydrophilic group is 1.7-4.0Å from the side chain atom;
  - (vii) an R<sub>2</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD;
- 10 (viii) an R<sub>3</sub>'-substituent comprising a hydrophobic group that interacts with a side chain atom of a phenylalanine, glycine or methionine corresponding to a residue selected from the group consisting of Phe215, Gly290, and Met388 of human TR-α, and Phe269, Gly344, Met442 of human TR-β, and wherein the hydrophobic group is 1.7-4.0Å from the side chain atom:
- 15 (ix) an R<sub>4</sub>'-substituent comprising an hydrogen bond donor or acceptor group that interacts with a side chain carbon or nitrogen atom of a histidine corresponding to residue His381 of human TR-α, and His435 of human TR-β, and wherein the hydrogen bond donor or acceptor group is 1.7-4.0Å from the side chain atom;
- 20 (x) an R<sub>5</sub>'-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD; and
  - (xi) and  $R_6$ '-substituent comprising a hydrophobic or hydrophilic group that fits spacially into the TR LBD.
- 25 22. The method of claim 19, wherein said compound comprises:

(i) a cyclic carbon atom that interacts with a carbon and oxygen atom of a methionine residue corresponding to Met259 of human TR-α, and Met313 of human TR-β, wherein the cyclic carbon is about 3.0 to 4.0Å from the carbon and oxygen atom of the methionine;

- 5 (ii) a cyclic carbon atom that interacts with a carbon atom of a leucine residue corresponding to Leu276 of human TR-α, and Leu330 of human TR-β, wherein the cyclic carbon is about 3.0 to 4.0Å from the carbon atom of the leucine;
- (iii) a cyclic carbon atom that interacts with a carbon atom of a leucine residue corresponding to Leu292 of human TR-α, and Leu346 of human TR-β,
   10 wherein the cyclic carbon is about 3.0 to 4.0Å from the carbon atom of the leucine;
  - (iv) a R<sub>3</sub>-substituent comprising an atom that interacts with a carbon atom of an isoleucine residue corresponding to Ile221 of human TR-α, and Ile275 of human TR-β, wherein the R3-substituent atom is about 3.0 to 4.0Å from the carbon atom of the isoleucine;
- 15 (v) a R<sub>3</sub>'-substituent comprising an atom that interacts with an oxygen atom of a glycine residue corresponding to Gly290 of human TR-α, and Gly344 of human TR-β, wherein the R3'-substituent atom is about 3.0 to 4.0Å from the carbon atom of the glycine; and
- (vi) a R<sub>4</sub>'-substituent comprising an atom selected from the group 20 consisting of oxygen and carbon that interacts with (a) a carbon and nitrogen atom of a histidine residue corresponding to His381 of human TR-α, and His435 of human TR-β, wherein the R4'-substituent atom is about 2.0 to 4.0Å from the carbon atom of the histidine; and (b) a carbon atom of a phenylalanine residue corresponding to Phe401 of human TR-α, and human Phe455 of TR-β, wherein said atom is about 3.0 to 4.0Å from the carbon atom of the phenylalanine.

PCT/US98/25296 WO 99/26966

> 23. The method according to claim 21,

wherein R<sub>1</sub> is

-O-CH<sub>2</sub>CO<sub>2</sub>H, -NHCH<sub>2</sub>CO<sub>2</sub>H,

-CO<sub>2</sub>H, -CH<sub>2</sub>CO<sub>2</sub>H, -CH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>H, -CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub>H, 5

> -CH<sub>2</sub>CH[NHCOCHφ<sub>2</sub>]CO<sub>2</sub>H, -CH<sub>2</sub>CH(NH<sub>2</sub>)CO<sub>2</sub>H,

> -CH2CH[NH-FMOC]CO2H, -CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub> ]CO<sub>2</sub>H,

> -CH<sub>2</sub>CH[NH-tBOC]CO<sub>2</sub>H, or a carboxylate connected to the ring with a 0 to 3

carbon linker,

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-CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CHNH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub>,  $-PO_3H_2$ 

-CH<sub>2</sub>CH[NHCO(CH<sub>2</sub>)<sub>15</sub>CH<sub>3</sub>]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub>CH[NHCOCH $\phi_2$ ]PO<sub>3</sub>H<sub>2</sub>,

-CH<sub>2</sub>CH[NH-FMOC]PO<sub>3</sub>H<sub>2</sub>, -CH<sub>2</sub> CH[NH-tBOC]PO<sub>3</sub>H<sub>2</sub>, or a phosphate or

phosphonate connected to the ring with a 0 to 3 carbon linker,

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-CH<sub>2</sub>CH<sub>2</sub>SO<sub>3</sub>H, -CH<sub>2</sub>CHNH<sub>2</sub>SO<sub>3</sub>H, -SO<sub>3</sub>H, -CH<sub>2</sub>SO<sub>3</sub>H,

-CH2CH[NHCO(CH2)15CH3]SO3H, -CH<sub>2</sub>CH[NHCOCHφ<sub>2</sub>]SO<sub>3</sub>H,

-CH<sub>2</sub>CH[NH-FMOC]SO<sub>3</sub>H, -CH<sub>2</sub> CH[NH-tBOC]SO<sub>3</sub>H, or a sulfate or sulfite

connected to the ring with a 0 to 3 carbon linker,

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or acts as the functional equivalent of CH2CH(NH2)CO2H of T3 in the molecular recognition domain when bound to a TR, wherein said R1 can be optionally substituted with an amine,

25 wherein R<sub>2</sub> is

H, halogen, CF<sub>3</sub>, OH, NH<sub>2</sub>, SH, CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

## 5 wherein R<sub>3</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

### 10 wherein R<sub>5</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR, and  $R_3$  can be identical to  $R_5$ ,

# 15 wherein R<sub>6</sub> is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR, and R<sub>2</sub> can be identical to R<sub>6</sub>,

## 20 wherein R2' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -N<sub>3</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

wherein R<sub>3</sub>' is any hydrophobic group, including

halogen, -CF<sub>3</sub>, -SH, alkyl, aryl, 5- or 6-membered heterocycle, cyano, or acts as the functional equivalent of I in the molecular recognition domain when bound to a TR,

### 5 wherein R<sub>4</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, -SH, -CH<sub>3</sub>, -Et, or akyl, aryl or 5- or 6-membered heterocyclic aromatic attached through urea or carbamate linkages to O or N or S at the R<sub>4</sub>' position, or acts as the functional equivalent of OH in the molecular recognition domain when bound to a TR,

### wherein R5' is

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-H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is optionally connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups,

wherein R<sub>6</sub>' is

-H, halogen, -CF<sub>3</sub>, -OH, -NH<sub>2</sub>, -SH, -CH<sub>3</sub>, -Et, or acts as the functional equivalent of H in the molecular recognition domain when bound to a TR,

wherein X is

O, S, SO<sub>2</sub>, NH, NR<sub>7</sub>, CH<sub>2</sub>, CHR<sub>7</sub>, CR<sub>7</sub>R<sub>7</sub>, wherein R<sub>7</sub> is alkyl, aryl or 5- or 6-membered heterocyclic aromatic,

and wherein said TR LBD ligand has an apparent Kd for binding TR LBD of 1 TM or less.

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24. The method of claim 23, wherein

R<sub>1</sub> is carboxylate, phosphonate, phosphate or sulfite and is connected to the ring with a 0 to 3 carbon linker,

R<sub>2</sub> is H,

15 R<sub>3</sub> is -I, -Br, or -CH<sub>3</sub>,

 $R_5$  is -I, -Br, or -CH<sub>3</sub>,

R<sub>6</sub> is H,

R<sub>2</sub>' is H,

R<sub>3</sub>' is -I, -Br, -CH<sub>3</sub>, -iPr, -phenyl, benzyl, or 5- or 6-membered ring 20 heterocycles,

R<sub>4</sub>' is -OH, -NH<sub>2</sub>, and -SH,

R<sub>5</sub>' is -H, -OH, -NH<sub>2</sub>, -N(CH<sub>3</sub>)<sub>2</sub> -SH -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate, sulfate, branched or straight chain alkyl having 1 to 9 carbons, substituted or unsubstituted aryl, wherein said substituted aryl is substituted with halogen or 1 to 5 carbon alkyl and wherein said aryl is

optionally connected to the ring by a -CH<sub>2</sub>-, aromatic heterocycle having 5 to 6 atoms, wherein said heterocycle may be substituted with one or more groups selected from -OH, -NH<sub>2</sub>, -SH, -NH<sub>3</sub>, -N(CH<sub>3</sub>)<sub>3</sub>, carboxylate, phosphonate, phosphate or sulfate, heteroalkyl, arylalkyl, heteroaryl alkyl, polyaromatic, or polyheteroaromatic, wherein said R<sub>5</sub>' may be substituted with polar or charged groups, and

R<sub>6</sub>' is H.

5

- 25. The method of claim 19, wherein said compound fits spatially and 10 preferentially into TR LBD isoform  $\alpha$  (TR- $\alpha$ ).
- 26. The method of claim 25, wherein said compound comprises an anionic group that interacts with the side chain oxygen or carbon of a serine residue corresponding to Ser277 of human TR-α, and wherein the anionic group is 1.7-4.0Å
  15 from the side chain atom.
  - 27. The method of claim 19, wherein said compound fits spatially and preferentially into TR LBD isoform  $\beta$  (TR- $\beta$ ).
- 28. The method of claim 27, wherein said compound comprises an anionic group that interacts with the side chain nitrogen of an arginine corresponding to Asn331 of human TR-β, and the anionic group is 1.7-4.0Å from the side chain atom.
- 29. The method of claim 19, wherein said compound binds to a TR LBD 25 isoform with greater affinity than thyronine or triiodothyronine.

30. The method of claim 1, wherein said compound comprises a cyclic carbon atom that interacts with a carbon and oxygen atom of a methionine residue corresponding to Met259 of human TR-I, and Met313 of human TR-β, wherein the cyclic carbon is about 3.0 to 4.0Å from the carbon and oxygen atom of the methionine.

31. The method of claim 30, wherein said cyclic carbon is inner ring carbon C11.

10

32. The method of claim 1, wherein said compound comprises a cyclic carbon atom that interacts with a carbon atom of a leucine residue corresponding to Leu276 of human TR-α, and Leu330 of human TR-β, wherein the cyclic carbon is about 3.0 to 4.0Å from the carbon atom of the leucine.

15

- 33. The method of claim 32, wherein said cyclic carbon is selected from the group consisting of inner ring carbons C7 and C9.
- 34. The method of claim 1, wherein said compound comprises a cyclic 20 carbon atom that interacts with a carbon atom of a leucine residue corresponding to Leu292 of human TR-α, and Leu346 of human TR-β, wherein the cyclic carbon is about 3.0 to 4.0Å from the carbon atom of the leucine.
- 35. The method of claim 34, wherein said cyclic carbon is selected from25 the group consisting of outer ring carbons C6 and C8.

36. The method of claim 1, wherein said R<sub>3</sub>-substituent comprises an atom that interacts with a carbon atom of an isoleucine residue corresponding to Ile221 of human TR-α, and Ile275 of human TR-β, wherein the R<sub>3</sub>-substituent atom is about
5 3.0 to 4.0Å from the carbon atom of the isoleucine.

- 37. The method of claim 1, wherein said R<sub>3</sub>'-substituent comprises an atom that interacts with an oxygen atom of a glycine residue corresponding to Gly290 of human TR-α, and Gly344 of human TR-β, wherein the R<sub>3</sub>'-substituent atom is 10 about 3.0 to 4.0Å from the carbon atom of the glycine.
- 38. The method of claim 1, wherein said R<sub>4</sub>'-substituent comprises an atom selected from the group consisting of oxygen and carbon that interacts with a carbon and nitrogen atom of a histidine residue corresponding to His381 of human 15 TR-α, and His435 of human TR-β, wherein the R<sub>4</sub>'-substituent atom is about 2.0 to 4.0Å from the carbon atom of the histidine.
- 39. The method of claim 1, wherein said R<sub>4</sub>'-substituent comprises an oxygen atom that interacts with a carbon atom of a phenylalanine residue 20 corresponding to Phe401 of human TR-α, and human Phe455 of TR-β, wherein said atom is about 3.0 to 4.0Å from the carbon atom of the phenylalanine.
- 40. A method for identifying a thyroid hormone receptor (TR) agonist or antagonist ligand that selectively modulates the activity of a TR compared to other nuclear receptors, said method comprising the steps of:

providing the atomic coordinates of a TR ligand binding domain (TR LBD) to a computerized modeling system;

modeling ligands which fit spacially into the TR LBD and which interact with conformationally constrained residues of a TR LBD conserved among 5 TR isoforms; and

identifying in a biological assay for TR activity a ligand which selectively binds to said TR and increases or decreases the activity of said TR, whereby a TR agonist or antagonist that selectively modulates the activity of a TR is identified.

10

- 41. A peptide, peptidomatic or synthetic molecule identified by the method of any one of claims 19 or 40, with the proviso that said molecule is other than a thyronine or thyronine-like compound disclosed in a reference cited in Appendix I.
- 15 42. A machine-readable data storage medium, comprising a data storage material encoded with machine readable data which, when using a machine programmed with instructions for using said data, is capable of displaying a graphical three-dimensional representation of a molecule or molecular complex for a thyroid hormone ligand binding pocket comprising structure coordinates of TR-α amino acids 20 corresponding to human TR-α amino acids Met259, Leu276, and Ile221, or a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

43. A machine-readable data storage medium, comprising a data storage material encoded with machine readable data which, when using a machine programmed with instructions for using said data, is capable of displaying a graphical three-dimensional representation of a molecule or molecular complex for a thyroid 5 hormone ligand binding pocket comprising structure coordinates of TR-α amino acids corresponding to human TR-α amino acids Leu292, His381, Gly290 and Phe401, or a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

10

44. The machine-readable storage medium according to any one of claims 42 or 43, wherein said binding pocket comprises structure coordinates of TR- $\alpha$  amino acids corresponding to human TR- $\alpha$  amino acids Met259, Leu276, Leu292, His381, Gly290, Ile221 and Phe401.

15

- 45. The machine-readable storage medium according to claim 44. wherein said binding pocket comprises structure coordinates of TR-α amino acids corresponding to human TR-α amino acids Arg228, Arg262 and Arg266.
- 20 46. The machine-readable storage medium according to claim 44, wherein said binding pocket comprises structure coordinates of TR-α amino acids corresponding to human TR-α amino acids Ser260, Ala263 and Ile299.

47. The machine-readable storage medium according to claim 44, wherein said binding pocket comprises structure coordinates of TR- $\alpha$  amino acids corresponding to human TR- $\alpha$  amino acids Phe218, Ile221 and Ile222.

- 5 48. The machine-readable storage medium according to claim 44, wherein said binding pocket comprises structure coordinates of TR-α amino acids corresponding to human TR-α amino acids Phe215, Gly290 and Met388.
- 49. The machine-readable storage medium according to claim 44, wherein 10 said binding pocket comprises structure coordinates of a TR-α amino acid corresponding to human TR-α amino acid Ser277.
- 50. A machine-readable data storage medium, comprising a data storage material encoded with machine readable data which, when using a machine programmed with instructions for using said data, is capable of displaying a graphical three-dimensional representation of a molecule or molecular complex for a thyroid hormone ligand binding pocket comprising structure coordinates of TR-β amino acids corresponding to human TR-β amino acids Met313, Leu330, and Ile275, or a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.
- 51. A machine-readable data storage medium, comprising a data storage material encoded with machine readable data which, when using a machine programmed with instructions for using said data, is capable of displaying a graphical

three-dimensional representation of a molecule or molecular complex for a thyroid hormone ligand binding pocket comprising structure coordinates of TR-β amino acids corresponding to human TR-β amino acids Leu346, His435, Gly344, and Phe455, or a homologue of said molecule or molecular complex, wherein said homologue comprises a binding pocket that has a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

- 52. The machine-readable data storage medium according to any one of claims 50 or 51, wherein said binding pocket comprises structure coordinates of TR-β amino acids corresponding to human TR-β amino acids Met313, Leu330, Leu346, His435, Gly344, Ile275 and Phe455.
- 53. The machine-readable data storage medium according to claim 52,
   wherein said binding pocket comprises structure coordinates of TR-β amino acids
   15 corresponding to human TR-β amino acids Arg282, Arg316 and Arg320.
  - 54. The machine-readable data storage medium according to claim 52, wherein said binding pocket comprises structure coordinates of TR-β amino acids corresponding to human TR-β amino acids Ser314, Ala317 and Ile352.

20

55. The machine-readable data storage medium according to claim 52, wherein said binding pocket comprises structure coordinates of TR-β amino acids corresponding to human TR-β amino acids Phe272, Ile275 and Ile276.

56. The machine-readable data storage medium according to claim 52, wherein said binding pocket further comprises structure coordinates of TR-β amino acids corresponding to human TR-β amino acids Phe269, Gly344 and Met442.

- 5 57. The machine-readable data storage medium according to claim 52, wherein said binding pocket comprises structure coordinates of a TR-β amino acid corresponding to human TR-β amino acid Asn331.
- 58. The machine-readable data storage medium according to claim 52, 10 wherein said molecule or molecular complex is defined by the set of structure coordinates selected from the group consisting coordinates depicted in Appendix 3, 4, 5 and 6, or a homologue of said molecule or molecular complex, said homologue having a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.

15

- 59. The machine-readable data storage medium according to claim 52, wherein said molecule or molecular complex is defined by the set of structure coordinates selected from the group consisting coordinates depicted in Appendix 7 and 8, or a homologue of said molecule or molecular complex, said homologue 20 having a root mean square deviation from the backbone atoms of said amino acids of not more than 1.5Å.
- 60. A machine-readable data storage medium comprising a data storage material encoded with a first set of machine readable data which, when combined 25 with a second set of machine readable data, using a machine programmed with

instructions for using said first set of data and said second set of data, can determine at least a portion of the structure coordinates corresponding to the second set of machine readable data, wherein: said first set of data comprises a Fourier transform of at least a portion of the structural coordinates selected from the group consisting of coordinates depicted in Appendix 3, 4, 5, 6, 7 and 8; and said second set of data comprises an X-ray diffraction pattern of a molecule or molecular complex.

WO 99/26966

PCT/US98/25296

### APPENDIX 1

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WO 99/26966

PCT/US98/25296

# APPENDIX 2

# Table 8

Dimit	Amino Acid	Amino Acid	Distance
Atom	in full length α	Atom	Å
C16	215-PHE	CD1	3.98
C16	215-PHE	CE1	3.86
C19	218-PHE	0	3.69
C16	218-PHE	СВ	3.89
C18	218-PHE	СВ	3.92
C19	218-PHE	СВ	4.13
C18	218-PHE	CD2	3.77
C16	219-THR	CG2	3.68
C19	221-ILE	CG1	4.11
C6	222-ILE	CD1	4.18
C8	222-ILE	CD1	3.72
C10	222-ILE	CD1	3.53
C12	222-ILE	CD1	3.85
01	222-ILE	CD1	4.13
C13	225-ALA	C8	3.64
04	225-ALA	C8	4.02
04	228-ARG	CZ	3.96
C17	228-ARG	NH2	3.36
O3	228-ARG	NH2	3.58
04	228-ARG	NH2	2.86
C10	256-MET	SD	3.70
C12	256-MET	SD	3.89
C10	256-MET	CE	3.88
C12	256-MET	CE	3.83
C11	259-MET	С	4.03
C11	259-MET	0	3.66
C15	259-MET	0	3.42
N1	259-MET	0	3.71
C1	259-MET	C8	4.20
C11	259-MET	C8	3.87
C13	259-MET	C8	4.09
C15	262-ARG	C8	4.03
C17	262-ARG	C8	3.58
O3	262-ARG	C8	3.62
04	262-ARG	C8	3.85
C17	262-ARG	CD	4.10
O4	262-ARG	CD	3.61
N1	263-ALA	N	3.71
C17	263-ALA	CA	3.69
NI	263-ALA	CB	3.46
O3	266-ARG	NHI	3.93

Dimit	Amino Acid	Amino Acid	Distance
Atom	in full length α	Atom	Å
NI	275-THR	0	3.62
N1	276-LEU	CA	3.51
Ni	276-LEU	C	3.92
C5	276-LEU	CD1	4.05
C19	276-LEU	CD1	4.04
C7	276-LEU	CD2	4.09
C9	276-LEU	CD2	3.95
C11	276-LEU	CD2	4.13
N1	276-LEU	CD2	4.17
C13	277-SER	N	4.14
C15	277-SER	N	3.79
C17	277-SER	N	3.69
NI	277-SER	N	3.30
O3	277-SER	N	3.19
C17	277-SER	CA	3.92
O3	277-SER	CA	3.35
C13	277-SER	OG	3.92
C7	287-LEU	CD2	3.90
C18	290-GLY	C	4.04
C18	290-GLY	0	3.54
C18	291-GLY	CA	4.04
C18	292-LEU	N	4.20
C2	292-LEU	CG	4.18
C4	292-LEU	CG	3.86
C6	292-LEU	CG	4.01
C2	292-LEU	CD1	3.88
C4	292-LEU	CD1	4.02
O2	292-LEU	CD1	4.07
C4	292-LEU	CD2	4.05
C6	292-LEU	CD2	3.72
C8	292-LEU	CD2	3.69
C10	292-LEU	CD2	3.98
O1	292-LEU	CD2	4.16
C20	299-ILE	CD1	3.87
C8	381-HIS	CD2	3.90
C10	381-HIS	CD2	3.84
O1	381-HIS	GO2	3.40
01	381-HIS	CE1	3.72
C8	381-HIS	NE2	3.47
C10	381-HIS	NE2	3.51
01	381-HIS	NE2	2.64
C6	388-MET	CE	3.90
C8	401-PHE	CE1	4.19
O1	401-PHE	CE1	3.37

WO 99/26966

PCT/US98/25296

Dimit Atom	Amino Acid in full length α	Amino Acid Atom	<u>Distance</u> Å
C16	401-PHE	CZ	3.97
O1	401-PHE	CZ	3.28
NI	502-H <sub>2</sub> O	01	3.35
O3	502-H <sub>2</sub> O	O1	2.56
O3	503-H <sub>2</sub> O	O1	3.13
04	503-H <sub>2</sub> O	01	3.72
O4	504-H <sub>2</sub> O	O1	2.72

Legend to Table 8. The table lists the interactions with Dimit (DMT). The column headings are as follows:

- #1 The atom of Dimit that interacts with the amino acid of the receptor. These are also 5 numbered in Figure 32.
  - #2 The amino acid in the full length rTR $\alpha$  that interacts with the ligand.
  - #3 The name of the atom in the amino acid (standard nomenclature) where the interaction occurs.
  - #4 The distance in Å between Dimit and the protein atom.

WO 99/26966

PCT/US98/25296

Table 9

Triac	Amino Acid	Amino Acid	
Atom	in full length α	Atom	Distance Å
I1	218-PHE	0	3.52
11	221-ILE	CD1	4.16
I1	221-ILE	CG1	3.92
11	222-ILE	CA	4.15
Il	222-ILE	CB	4.03
Il	222-ILE	CG1	3.92
C8	222-ILE	CD1	4.12
C10	222-ILE	CD1	3.77
C12	222-ILE	CD1	3.79
C13	225-ALA	СВ	4.17
C3	225-ALA	CB	3.86
C10	256-MET	SD	3.45
C12	256-MET	SD	3.73
C10	256-MET	CE	3.66
C12	256-MET	CE	3.77
I3	256-MET	CE	3.89
C1	259-MET	O	3.93
C11	259-MET	0	3.24
O3	259-MET	0	4.09
C1	259-MET	СВ	3.89
C13	259-MET	O	3.74
C14	259-MET	0	3.96
Cl	259-MET	СВ	3.89
C11	259-MET	CB	3.68
C13	259-MET	СВ	4.01
C11	259-MET	CA	4.13
C13	259-MET	CA	4.20
<u>I3</u>	260-SER	CA	4.10
I3	260-SER	OG	4.19
C14	262-ARG	СВ	4.07
04	262-ARG	СВ	3.60
O3	263-ALA	N	3.79
C14	263-ALA	N	4.12
O3	263-ALA	CA	3.67
O3	263-ALA	CB	3.49
C11	263-ALA	CB	4.00
C14	266-ARG	CZ	3.89
03	266-ARG	CZ	4.01
04	266-ARG	CZ	3.03
C14	266-ARG	NHI	3.25
03	266-ARG	NH1	3.00
04	266-ARG	NH1	2.82

Triac	Amino Acid	Amino Acid	
Atom	in full length α	Atom	Distance Å
C14	266-ARG	NH2	3.48
03	266-ARG	NH2	4.01
04	266-ARG	NH2	2.34
03	275-THR	С	4.02
C14	275-THR	0	4.20
O3	275-THR	0	3.20
O3	278-LEU	CA	3.11
03	276-LEU	C	3.52
O3	276-LEU	N	4.04
C14	276-LEU	CA	3.98
O3	276-LEU	CA	3.11
C14	276-LEU	C	3.98
O3	276-LEU	СВ	3.95
O2	276-LEU	CD1	4.03
I1	276-LEU	CD1	4.10
C7	276-LEU	CD2	3.84
C9	276-LEU	CD2	3.73
CII	276-LEU	CD2	4.06
O2	276-LEU	CD2	4.10
03	276-LEU	CD2	3.91
C13	277-SER	N	4.06
C14	277-SER	N	3.13
O4	277-SER	N	3.28
O3	277-SER	N	3.05
C14	277-SER	CA	3.76
O4	277-SER	CA	3.52
C3	277-SER	OG	3.87
C13	277-SER	OG	4.02
C14	277-SER	OG	4.14
I2	290-GLY	0	3.57
I2	292-LEU	CG	3.94
C4	292-LEU	CG	3.95
C6	292-LEU	CG	3.65
C8	292-LEU	CG	4.02
C2	292-LEU	CD1	4.11
C4	292-LEU	CD1	3.85
C6	292-LEU	CD1	4.02
I2	292-LEU	CD2	3.98
C4	292-LEU	CD2	4.11
C6	292-LEU	CD2	3.44
C8	292-LEU	CD2	3.28
C10	292-LEU	CD2	3.88
01	292-LEU	CD2	3.35
	299-ILE	CD1	3.77

Triac	Amino Acid	Amino Acid	
Atom	in full length α	Atom	Distance Å
C8	381-HIS	CD2	3.87
C10	381-HIS	CD2	3.90
O1	381-HIS	GO2	3.20
O1	381-HIS	CEI	3.82
C8	381-HIS	NE2	3.57
C10	381-HIS	NE2	3.52
Ol	381-HIS	NE2	2.64
O1	388-MET	CE	4.03
01	401-PHE	CE1	3.86
01	401-PHE	CZ	3.70
C13	460-H <sub>2</sub> 0	01	4.00

Legend to Table 9. The table lists the interactions with Triac. The column headings are as follows:

- #1 The atom of Triac that interacts with the amino acid of the receptor. These are also 5 numbered in Figure 32.
  - #2 The amino acid in the full length  $rTR\alpha$  that interacts with the ligand.
  - #3 The name of the atom in the amino acid (standard nomenclature) where the interaction occurs.
  - #4 The distance in Å between Triac and the protein atom.

Table 10

IpBR <sub>2</sub> Atom	Amino Acid	Amino Acid	Distance
1pD102110000	in full length α	Atom	Å
C16	215-PHE	CD1	4.01
C16	215-PHE	CEI	3.78
BR1	218-PHE	0	3.24
BR1	218-PHE	C	3.98
C16	218-PHE	СВ	3.81
C18	218-PHE	СВ	3.92
BR1	218-PHE	СВ	4.08
C18	218-PHE	CD2	3.92
C16	219-THR	CG2	3.45
BR1	221-ILE	CG1	3.81
BR1	221-ILE	CD1	4.07
BR1	222-ILE	СВ	3.81
BR1	222-ILE	CG1	3.97
C6	222-ILE	CD1	4.07
C8	222-ILE	CD1	3.64
C10	222-ILE	CD1	3.50
C12	222-ILE	CD1	3.82
01	222-ILE	CD1	4.08
C13	225-ALA	СВ	3.76
04	225-ALA	CB	4.01
04	228-ARG	CZ	3.92
C17	228-ARG	NH2	3.26
03	228-ARG	NH2	3.43
04	228-ARG	NH2	2.79
C10	256-MET	SD	3.65
C12	256-MET	SD	3.71
C10	256-MET	CE	3.90
C12	256-MET	CE	3.75
BR2	256-MET	CE	4.03
C11	259-MET	C	3.98
C11	259-MET	0	3.52
C15	259-MET	0	3.44
N1	259-MET	0	3.76
C11	259-MET	CB	3.87
N1	262-ARG	С	4.03
C15	262-ARG	CB	4.03
C17	262-ARG	СВ	3.56
03	262-ARG	CB	3.55
04	262-ARG	СВ	3.91
C17	262-ARG	CD	4.09
04	262-ARG	CD	3.71
N1	263-ALA	N	3.61

IpBR <sub>2</sub> Atom	Amino Acid	Amino Acid	<u>Distance</u>
	in full length α	Atom .	Ä
N1	263-ALA	CA	3.59
N1	263-ALA	СВ	3.54
03	266-ARG	NH1	3.93
N1	275-THR	0	3.43
NI	276-LEU	CA	3.46
N1	276-LEU	С	3.83
C5	276-LEU	CD1	4.02
C7	276-LEU	CD2	4.00
C9	276-LEU	CD2	3.81
C11	276-LEU	CD2	3.91
C13	277-SER	N	3.79
C15	277-SER	N	3.63
C17	277-SER	N	3.70
N1	277-SER	N	3.17
03	277-SER	N	3.37
C17	277-SER	CA	3.89
03	277-SER	CA	3.43
C13	277-SER	OG	3.66
02	287-LEU	CD1	4.05
C18	290-GLY	C	4.04
C18	290-GLY	0	3.48
C18	291-GLY	CA	4.02
C4	292-LEU	CG	3.89
C6	292-LEU	CG	4.02
C2	292-LEU	CD1	3.79
C4	292-LEU	CD1	3.96
02	292-LEU	CD1	3.97
C4	292-LEU	CD2	4.07
C6	292-LEU	CD2	3.75
C8	292-LEU	CD2	3.67
C10	292-LEU	CD2	3.92
BR2	299-ILE	CD1	3.68
C8	381-HIS	CD2	3.92
C10	381-HIS	CD2	3.78
01	381-HIS	GD2	3.50
01	381-HIS	CE1	3.62
C8	381-HIS	NE2	3.36
C10	381-HIS	NE2	3.34
01	381-HIS	NE2	2.62
C8	401-PHE	CE1	4.02
01	401-PHE	CE1	3.19
C16	401-PHE	CZ	4.03
01	401-PHE	CZ	3.06
03	502-H <sub>2</sub> O	01	3.40

10

WO 99/26966 PCT/US98/25296

IpBR <sub>2</sub> Atom	Amino Acid	Amino Acid	Distance
	in full length α	Atom	Å
Nl	502-H20	01	3.12
04	503-H <sub>2</sub> O	01	3.20
C17	503-H20	01	3.04
03	503-H <sub>2</sub> O	01	2.27
C15	504-H20	01	4.01
C17	504-H <sub>2</sub> O	01	2.99
03	504-H2O	01	3.80
04	504-H <sub>2</sub> O	01	1.78

Legend to Table 10. The table lists the interactions with IpBr2. The column headings are as follows:

- #1 The atom of IpBr2 that interacts with the amino acid of the receptor. These are also 5 numbered in Figure 32.
  - #2 The amino acid in the full length rTR $\alpha$  that interacts with the ligand.
  - #3 The name of the atom in the amino acid (standard nomenclature) where the interaction occurs.
  - #4 The distance in Å between IpBr2 and the protein atom.

Table 11

I abic		
Amino Acid	Amino Acid	Distance
in full length α	Atom	Å
215-PHE		4.08
218-PHE	0	3.19
218-PHE		3.99
218-PHE	CB	4.04
218-PHE		3.79
218-PHE	CB	3.99
221-ILE	CG1	4.01
222-ILE	СВ	3.95
222-ILE	CG1	3.91
222-ILE	CD1	3.99
222-ILE	CD1	3.57
222-ILE	CD1	3.68
225-ALA	СВ	3.66
	СВ	4.04
228-ARG	NH1	3.23
228-ARG	CZ	3.45
	NH2	3.54
228-ARG	NH2	3.90
228-ARG	NH2	2.86
256-MET	SD	3.73
256-MET	SD	3.90
1	CE	3.97
	CE	3.92
	CE	3.89
259-MET	С	3.95
	0	3.59
259-MET	0	3.51
259-MET	0	3.88
259-MET	СВ	4.06
259-MET	СВ	3.77
259-MET	CB	3.96
	СВ	3.61
	СВ	4.02
	СВ	3.65
262-ARG	СВ	3.92
262-ARG	CD	3.72
		3.81
		3.81
		3.63
		3.54
		3.38
	C	3.73
		4.00
	Amino Acid in full length α  215-PHE  218-PHE  218-PHE  218-PHE  218-PHE  218-PHE  221-ILE  222-ILE  222-ILE  222-ILE  222-ILE  222-ILE  222-ILE  225-ALA  225-ALA  228-ARG  228-ARG  228-ARG  228-ARG  228-ARG  228-ARG  256-MET  256-MET  256-MET  256-MET  259-MET  259-MET	in full length α         Atom           215-PHE         CD1           218-PHE         0           218-PHE         CB           218-PHE         CB           218-PHE         CB           221-ILE         CB           221-ILE         CB           222-ILE         CD1           222-ILE         CD1           222-ILE         CD1           222-ILE         CD1           225-ALA         CB           225-ALA         CB           228-ARG         NH1           228-ARG         NH2           259-ME

WO 99/26966

# PCT/US98/25296

T3 Atom	Amino Acid	Amino Acid	Distance
***************************************	in full length α	Atom	Å
C7	276-LEU	CD1	4.05
02	276-LEU	CD1	4.03
C7	276-LEU	CD2	3.80
<u>C9</u>	276-LEU	CD2	3.70
CII	276-LEU	CD2	4.01
C14	277-SER	N	3.67
C15	277-SER	N	3.62
N1	277-SER	N	3.07
03	277-SER	N	3.24
C15	277-SER	CA	3.77
03	277-SER	CA	3.34
C13	277-SER	OG	3.92
12	290-GLY	0	3.50
C4	292-LEU	CG	3.95
C8	292-LEU	CG	3.83
C2	292-LEU	CD1	4.07
C4	292-LEU	CD1	3.99
C4	292-LEU	CD2	4.09
C6	292-LEU	CD2	3.58
C8	292-LEU	CD2	3.50
C10	292-LEU	CD2	3.96
01	292-LEU	CD2	3.71
13	299-ILE	CD1	3.74
C8	381-HIS	CD2	3.94
C10	381-HIS	CD2	3.97
01	381-HIS	CD2	3.39
01	381-HIS	CD1	3.82
C8	381-HIS	NE2	3.47
C10	381-HIS	NE2	3.55
01	381-HIS	NE2	2.70
01	388-MET	CE	3.88
01	401-PHE	CEI	3.52
01	401-PHE	CZ	3.32
C14	502-H20	O1	4.01
C15	502-H2O	O1	3.61
03	502-H20	01	2.51
C15	503-H2O	O1	3.31
04	503-H <sub>2</sub> O	01	3.10
N1	502-H <sub>2</sub> O	01	3.27
03	503-H2O	01	2.81
C15	504-H2O	Oi	3.92
04	504-H2O	01	2.73
Total Table 11	The table lists the inter		he column he

Legend to Table 11. The table lists the interactions with T3. The column headings are as follows:

#1 The atom of T3 that interacts with the amino acid of the receptor. These are also numbered in Figure 32.

#2 The amino acid in the full length rTRα that interacts with the ligand.

#3 The name of the atom in the amino acid (standard nomenclature) where the 5 interaction occurs.

#4 The distance in Å between T3 and the protein atom.

Table 12

Triac	Amino Acid	Amino Acid	•
Atom	in full length hTR β	Atom	Distance Å
I2	269-PHE	CD1	3.75
12	269-PHE	CE1	3.88
I1	272-PHE	С	4.03
Il	272-PHE	0	3.54
Ĭ1	275-ILE	CG1	3.93
11	276-ILE	CG1	4.02
C3	279-ALA	СВ	3.81
C13	279-ALA	СВ	3.87
C10	310-MET	SD	3.72
C12	310-MET	SD	3.78
C10	310-MET	CE	4.02
C12	310-MET	CE	3.92
I3	310-MET	CE	3.93
C13	313-MET	CA	3.94
C11	313-MET	C	3.72
C1	313-MET	0	3.79
C11	313-MET	0	3.12
C13	313-MET	0	3.55
Cl	313-MET	СВ	4.00
C11	313-MET	СВ	3.82
C13	313-MET	СВ	3.76
C13	313-MET	CG	3.88
O3	316-ARG	CB	3.99
O4	317-ALA	CA	4.08
04	317-ALA	CA	4.10
C11	317-ALA	СВ	3.70
I3	317-ALA	CB	4.10
O4	317-ALA	СВ	4.06
O4	320-ARG	NHI	3.58
03	320-ARG	NH2	3.55
04	320-ARG	NH2	4.04
04	329-THR	0	3.55
04	330-LEU	CA	3.42
04	330-LEU	C	3.77
C3	330-LEU	СВ	4.06
C5	330-LEU	CB	4.08
<u>C1</u>	330-LEU	CD2	4.07
C3	330-LEU	CD2	4.00
C5	330-LEU	CD2	3.73
C7	330-LEU	CD2	3.51
C9	330-LEU	CD2	3.54
CII	330-LEU	CD2	3.86

Triac	Amino Acid	Amino Acid	
Atom	in full length hTR β	Atom	Distance Å
C15	331-ASN	N	3.55
O3	331-ASN	N	3.74
O4	331-ASN	N	3.12
03	331-ASN	CA	4.02
I2	344-GLY	0	3.87
C6	346-LEU	CD2	3.87
C8	346-LEU	CD2	3.84
01	346-LEU	CD2	3.91
13	353-ILE	CD1	3.51
C8	435-HIS	CD2	3.93
C10	435-HIS	CD2	3.79
01	435-HIS	CD2	3.33
01	435-HIS	CEI	3.81
C8	435-HIS	NE2	3.42
C10	435-HIS	NE2	3.33
01	435-HIS	NE2	2.67
01	442-MET	SD	3.96
01	442-MET	CE	3.72
12	442-MET	SD	4.01
O1	455-PHE	CE1	3.92
01	455-PHE	CZ	3.50

Legend to Table 12. The table lists the interactions with Triac. The column headings are as follows:

<sup>#1</sup> The atom of Triac that interacts with the amino acid of the receptor. These are also 5 numbered in Figure 32.

<sup>#2</sup> The amino acid in the full length hTRβ that interacts with the ligand.

<sup>#3</sup> The name of the atom in the amino acid (standard nomenclature) where the interaction occurs.

<sup>#4</sup> The distance in Å between Triac and the protein atom.

WO 99/26966

# PCT/US98/25296

Table 13

GC1	Amino Acid	Amino Acid	
Atom	in full length TR β	Atom	Distance Å
C16	269-PHE	CE1	3.99
C19	272-PHE	0	3.85
C16	272-PHE	CB	3.98
C16	273-THR	CG2	3.76
C19	275-ILE	CG1	3.98
C19	276-ILE	CA	3.98
C2	276-ILE	CD1	3.88
C8	276-ILE	CD1	3.77
C10	276-ILE	CD1	3.58
C12	276-ILE	CD1	3.62
C19	276-ILE	CD1	3.56
C1	279-ALA	СВ	3.68
C3	279-ALA	СВ	3.56
O5	279-ALA	CB	3.11
O4	279-ALA	СВ	3.90
O3	282-ARG	CZ	3.53
C17	282-ARG	NH1	3.87
O3	282-ARG	NH1	3.20
O4	282-ARG	NH1	3.85
C17	282-ARG	NH2	3.63
O3	282-ARG	NH2	3.00
C10	310-MET	SD	3.86
C12	310-MET	SD	3.91
C11	313-MET	C	3.85
C11	313-MET	0	3.41
C15	313-MET	0	3.87
C20	313-MET	0	3.99
C11	313-MET	СВ	3.79
C1	313-MET	CG	3.94
C11	313-MET	CG	3.91
O5	313-MET	CG	3.87
O4	313-MET	CG	3.79
C20	314-SER	CA	4.00
C17	316-ARG	CB	3.95
C17	316-ARG	CD	3.80
O3	316-ARG	CD	3.83
O4	316-ARG	CD	3.51
C20	317-ALA	CB	3.93
C7	330-LEU	CD2	3.56
C9	330-LEU	CD2	3.63
C21	330-LEU	CD2	3.90
O5	331-ASN	N	3.62

WO 99/26966

PCT/US98/25296

GC1	Amino Acid	Amino Acid	
Atom	in full length TR β	Atom	Distance Å
C15	331-ASN	N	3.67
C18	344-GLY	0	3.60
C18	346-LEU	CG	3.89
C6	346-LEU	CD2	3.77
C8	346-LEÚ	CD2	3.80
C10	435-HIS	CD2	3.89
01	435-HIS	CD2	3.64
O1	435-HIS	CE1	3.79
C8	435-HIS	NE2	3.44
C10	435-HIS	NE2	3.33
O1	435-HIS	NE2	2.77
01	455-PHE	CE1	3.40
O1	455-PHE	CZ	3.22

Legend to Table 13. The table lists the interactions with GC1. The column headings are as follows:

- #1 The atom of GC1 that interacts with the amino acid of the receptor. These are also 5 numbered in Figure 32.
  - #2 The amino acid in the full length hTR $\beta$  that interacts with the ligand.
  - #3 The name of the atom in the amino acid (standard nomenclature) where the interaction occurs.
  - #4 The distance in Å between GC1 and the protein atom.

WO 99/26966

PCT/US98/25296

Table 14
Coordination Structure of TR-α and Dimit

Structure         CHr-CH(NH <sub>3</sub> )(CO)H         -H         -CH <sub>3</sub> -CH <sub>3</sub> -H		Coo	rdina	ation S	tructu		'R-α	and Din	rit			
Structure	Coordina-	R <sub>1</sub>	R <sub>2</sub>	R <sub>3</sub>	R <sub>5</sub>	R <sub>6</sub>	R'2	R'3	R'4	R's	R'6	Х
CH <sub>1</sub> -CH(NH <sub>3</sub> )(CO <sub>3</sub> )H	ion		_									
AAA	tructure											
AA		-CH2-CH(NH2)(CO2)H	-H	-CH <sub>3</sub>	-CH <sub>3</sub>	-H	-H		-OH	-H	-H	0
SS	IA											
AA	S							Н3				
SS - H3				218		1		218				
AA	S	-		H3				H3				
SS         H3         H3				T	i			219				
AAA         221         H3         222         223         226	S							H3				
SS         H3         222         222         222         222         22 <td< td=""><td></td><td></td><td></td><td>221</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<>				221								
AA	SS			H3								
SS         H3         H4         H5-H6         H5-H6         H5-H6         H5-H6         H5-H6         H5-H6         H5-H6         H5-H6         H6         H5-H6         H6			-					222	222	222	222	
AA	SS							H3	H3	H3	Н3	
SS         H3         3		225										
AA       228       38       38       38       38       38       388       388       388       388       388       388       388       388       388       388       388       388       35       38       388	SS					1						
SS         H3         256         2:56         2:56         2:58         2:56         2:58         2:56         2:56         2:58         2:59         3:58         1:5-16						1						
AA	SS				1							
SS         H5-H6         H5		<del></del>		1		<b>1</b>					256	
AA       259       259         SS       H5-H6       H5-H6         AA       262       3         SS       H5-H6       3         AA       263       3         SS       H5-H6       3         AA       266       3         SS       loop       3         AA       275       3         SS       S3       S3         AA       276       276       276         SS       S3       S3       S3         AA       277       3       3         SS       loop       3       3         AA       290-291       3         SS       loop loop loop loop       3         AA       299       3         SS       H8       381         AA       381       381         SS       H11       H11	SS			1	1					H5-H6	H5-H6	
SS         H5-H6         H5-H6           AA         262         3           SS         H5-H6         3           AA         263         3           SS         H5-H6         3           AA         266         3           SS         loop         3           AA         275         3           SS         S3         S3           AA         276         276         276           SS         S3         S3         S3           AA         277         3         3           SS         loop         3           AA         290-291         3           SS         loop         1           AA         292         292         292           SS         loop loop loop loop         1           AA         299         3         381           SS         H11         H11           AA         388         1           SS         H11         H11		259		†		259						
AA 262 SS H5-H6 AA 263 SS H5-H6 AA 266 SS loop AA 275 SS S3 AA 276 SS S3 AA 277 SS loop AA 3275 SS S3 S	SS	H5-H6		1		H5-H6	1					
SS         H5-H6           AA         263           SS         H5-H6           AA         266           SS         loop           AA         275           SS         S3           AA         276         276         276           SS         S3         S3         S3           AA         277         SS         loop           AA         290-291         SS           AA         292         292         292           SS         loop loop loop loop           AA         299         SS           AA         381         381           SS         H11         H11           AA         388         SS		262										
AA       263         SS       H5-H6         AA       266         SS       loop         AA       275         SS       S3         AA       276       276       276         SS       S3       S3       S3         AA       277       SS       loop         AA       290-291       SS         AA       292       292       292         SS       loop loop loop loop         SS       H8       381       381         SS       H11       H11       H11	SS	H5-H6				1	1					
SS         H5-H6           AA         266           SS         loop           AA         275           SS         S3           AA         276         276         276           SS         S3         S3         S3           AA         277         SS         loop           AA         290-291         SS           SS         loop         SS           AA         292         292         292           SS         loop loop loop         SS           AA         299         SS         SS           AA         381         381           SS         H11         H11           AA         388         SS												
AA 266 SS loop AA 275 SS S3 AA 276 276 276 276 SS S3 S3 S3 S3 AA 277 SS loop AA 277 SS loop AA 290-291 SS loop AA 292 292 292 292 SS AA 299 SS AA 381 SS H8 SS H8 SS H111 H11 AA 388 SS H111 H11	SS	H5-H6			1							
SS         loop		266										Ī
SS         S3           AA         276         276         276         276           SS         S3         S2         S2         S2         S2         S2         S2         S2         S2         S2         S	SS	loop			1				1			
AA       276       276       276       276         SS       S3       S3       S3       S3         AA       277       270       270       270         SS       1000       10	AA	275		1	<b>†</b>							
SS         S3         S4         S4<	SS		**					T		Τ-	Ι	
SS         S3         S4         S4<		276			276							
SS         loop           AA         290-291           SS         loop           AA         292         292         292           SS         loop loop loop         loop           AA         299         SS         AA           AA         381         381           SS         HII         HII         HII           AA         388         HIII         HIII	SS	S3		S3	S3	S3						
AA	AA	277							Ĭ			
SS         loop         292         292         292         292         292         292         292         292         292         293 <td>SS</td> <td>loop</td> <td></td>	SS	loop										
AA     292     292     292     292       SS     loop     loop     loop     loop       AA     299						1	Ī					<u></u>
SS         loop         loop         loop         loop           AA         299	SS											
AA 299 SS H8 381 381 SS H11 H11 AA 388 SS H11 H11	AA						292		292		<u></u>	292
SS     H8       AA     381 381       SS     H11 H11       AA     388       SS     H11 H11	SS						loop	loop	loop	loop		loop
SS     H8       AA     381 381       SS     H11 H11       AA     388       SS     H11 H11	AA								ļ			
SS H11 H11	SS				H8							
SS	AA										<u> </u>	ļ
SS Hil	SS								HII	HII		1
1 <sup>7</sup>												
	SS		1									
	AA		1	7			Τ'''''	401	401			
SS H12 H12							1 "	H12	H12			
AA HOH502/HOH503 /HOH504												
SS	SS		<del>                                     </del>		1		1		1	T	1	1

AA = Amino Acid

SS = Secondary Structure

Coordination	R <sub>1</sub>	R <sub>2</sub>	R <sub>3</sub>	R <sub>5</sub>	R <sub>6</sub>	R'2	R'3	R'4	R's	R'6	Х
Structure			<u> </u>			ļ <u>.                                </u>		011	<del> </del>	<u> </u>	
	-CH₂-COOH	-H	-I	-1	-H	-H	-I	-OH	-H	-H	0
AA			218	<u> </u>		1			<u> </u>		
SS			H3					ļ			
AA			221			<u> </u>				<u> </u>	
SS			H3	L							ļ
AA			l		ļ		222	222	222	222	<u> </u>
SS							H3	H3	Н3	Н3	<u> </u>
AA	225					<u> </u>					<b></b>
SS	H3										<u> </u>
AA				256				ļ	256	256	<u> </u>
SS				H5-H6			1	<u> </u>	H5-H6	H5-H6	
AA	259				259			<u> </u>		<u> </u>	
SS	H5-H6		T		H5-H6		<u> </u>				
AA	262									<u> </u>	
SS	H5-H6										
AA	263										<u> </u>
SS	H5-H6										<u> </u>
AA	266										
SS	loop										
AA	275					]					
SS	S3										
AA	276		276	276	276						<u> </u>
SS	S3		S3	S3	S3	_					
AA	277							<u> </u>			
SS	loop										
AA							290			1	
SS							loop				<b></b> _
AA		1				292	292	292	292		292
SS	7		T			loop	loop	loop	loop		loop
AA				299							
SS				H8							
AA								381	381		
SS								H11	H11		
AA		1						388			
SS		T						H11			
AA		1					401	401			
SS		1					H12	H12			

5 AA = Amino Acid SS = Secondary Structure

Coordina-	R <sub>1</sub>	R <sub>2</sub>	R <sub>3</sub>	R <sub>5</sub>	R <sub>6</sub>	R'2	R'3	R'4	R's	R'6	X
tion											
Structure											
	-CH <sub>2</sub> -CH(NH <sub>2</sub> )(CO <sub>2</sub> )H	-H	-Br	-Br	-H	-H	-CH(CH <sub>3</sub> ) <sub>2</sub>	-OH	-H	-H	0
AA							215				
SS							Н3				<b></b>
AA			218				218				<b></b>
SS			H3				Н3				<u> </u>
AA							219				<b></b>
SS							Н3				<u> </u>
AA			221								<b></b>
SS			Н3								L
AA			11				222	222	222	222	<u> </u>
SS							H3	H3	Н3	Н3	ļ
AA	225										<del></del>
SS	Н3									ļ	Ļ
AA	228									<u> </u>	ļ
SS	H3		<b></b>					ļ	0.5.6		┞—
AA					256				256	256	<b></b>
SS					H5-H6	<u> </u>		ļ	H5-H6	Н5-Н6	├
AA	259				259	L		ļ		ļ	<b> </b>
SS	Н5-Н6		ļ		H5-H6	ļ	ļ	ļ		ļ	<b>ļ</b>
AA	262	<b></b>	<u> </u>					<b></b>	ļ	ļ	<del> </del>
SS	H5-H6		<b> </b>					ļ		<del> </del>	┼──
AA	263		ļ		ļ	ļ	ļ			<u> </u>	<b>↓</b>
SS	H5-H6	ļ						ļ			
AA	266		-				<u> </u>	-	<del> </del>		<del> </del>
SS	loop	ļ			<u> </u>	ļ	ļ	<del></del>		-	<del> </del>
AA	275		<del>  </del>			<u> </u>	<del></del>	<del> </del>	<del> </del>	<del></del>	┼
SS	S3		1000	076	200	<del> </del>	<b>_</b>	<del> </del>		<del> </del>	┼
AA	276	ļ	276	276	276	<del>↓</del> —		<del>├</del>		<del>                                     </del>	<del> </del>
SS	S3 277	ļ	S3	S3	S3	<del> </del>		<del> </del> -	<del> </del>	<del> </del> -	┼
AA SS	211	<del> </del>			ļ	<del> </del>		+		<del> </del>	+-
AA	<del> </del>	├			-	<del> </del>	290-291	<del> </del>	<del> </del>	1	+
SS		<del> </del>	<del></del>		<del> </del>	<del> </del>	loop	+	<del> </del>	<del> </del>	+
AA		1			<del> </del>	292	292	292	292	<del> </del>	292
SS	<del> </del>	-		<del> </del>	<del> </del>	loop	loop	loop	loop	<del> </del>	loop
AA		<del> </del>	+	299	<del> </del>	1000	1000	1.000	1.00	<del> </del>	1:55
SS	-	<del>                                     </del>	+	H8		+	+	+-	<del> </del>	1	+
AA	<del></del>	<del> </del>	+	110	<del> </del>	+	<del> </del>	381	381	<del> </del>	+
SS	-		+	-	+	<del> </del>	+	HII	H11	+	1
AA		+	+	<del>                                     </del>	+	<del> </del>	401	401	1	1	1
SS		┼	+	<del>                                     </del>		+	H12	H12	-	+	+ -
AA	HOH502/HOH503/	+		<del> </del>	<del> </del>	<del> </del>	1112	+****	<del>                                     </del>	+	+
SS	HOH504						ļ	<u> </u>			<del> </del>

AA = Amino Acid

SS = Secondary Structure

Table 17
Coordination Structure of TR-α and Dimit

Coordination	R <sub>1</sub>	R <sub>2</sub>	R <sub>3</sub>	R <sub>5</sub>	R <sub>6</sub>	R'2	R'3	R' <sub>4</sub>	R's	R'6	X
Structure											
	-CH <sub>2</sub> -CH(NH <sub>2</sub> )(CO <sub>1</sub> )H	-H	-1	-I	-H	-H	-1	-ОН	-Н	-H	0
AA				I			215				
SS							H3				
AA			218			218					
SS	-		H3			Н3					
AA			221								
SS			H3								
ĀĀ							222	222	222	222	
SS							H3	Н3	H3	Н3	
AA	225										
SS	H3										
AA	228										
SS	H3										
AA					256				256	256	
SS					H5-H6				H5-H6	H5-H6	
ĀĀ	259		1		259						L
SS	H5-H6		1		H5-H6						
ĀĀ	262				<u> </u>						
SS	H5-H6		<b>-</b>								<u> </u>
AA	263	1									
SS	H5-H6	<b>-</b>									<u> </u>
AA	275		1								<u> </u>
SS	S3			1						<u> </u>	
AA	276		276	276	276					<u> </u>	<u> </u>
SS	S3		S3	S3	S3						<u> </u>
AA	277										
SS											
AA							290				
SS							loop	<u> </u>			
AA				T		292	292	292	292		292
SS		1		1		loop	loop	loop	loop	<u> </u>	loop
AA				299				J		<u> </u>	
SS				H8							<u> </u>
AA								381	381	ļ	
SS								H11	HII	<b></b>	
AA								388		<u> </u>	
SS								H11	1		
AA				Τ			401	401			
SS		$\top$					H12	H12			
AA SS	HOH502/H0H503/ HOH504										

AA = Amino Acid SS = Secondary Structure

WO 99/26966

PCT/US98/25296

Table 18
Coordination Structure of TR-β and Triac

Coordination Structure	RI	R2	R3	R5	R6	R2'	R3	R4	R5	R6	Х
Structure	-CH <sub>2</sub> CO <sub>2</sub> H	Н	1	T	Н	Н	1	ОН	Н	Н	0
AA	0.120021.2						269				
SS							Н3				
AA			272		<del></del>						
SS			H3								
AA	<del></del>		275				i				
SS		-	H3								
AA			276								
SS			H3	<u>-</u>							
AA	279	279	-								
SS	H3	H3									
AA	1	· · · · · · · · · · · · · · · · · · ·	<del>                                     </del>	310					310	310	
SS			<del>                                     </del>	H5-H6					H5-H6	H5-H6	
AA	313		1		313						
SS	H5-H6				H5-H6						
AA	316										
SS	H5-H6										
AA	317				317		317				<u> </u>
SS	H5-H6				H5-H6		H5-H6				<u> </u>
AA	320										<u> </u>
SS	H5-H6					1					<u> </u>
AA	329				<u> </u>				<u> </u>		_
SS	S3			<u> </u>						<u> </u>	<u> </u>
AA	330	330	330	330	330				ļ	<u> </u>	<del> </del>
SS	S3	S3	S3	S3	S3			L		ļ	<u> </u>
AA	331		<u> </u>			<u> </u>		ļ			-
SS	loop	<u> </u>	<del></del>	<u> </u>	<u> </u>	<del>                                     </del>	-	<u> </u>	<del> </del>	<b>}</b>	₩
AA		<del> </del>		ļ <u>.</u>	ļ	-	344	<b> </b>	<del> </del>	<del> </del>	+
SS		J		<u> </u>	<u> </u>		loop	346	<del> </del>	<del> </del>	-
AA		<del> </del>	1	ļ		—	346	346		<del> </del>	+
SS		1		252	-	<del> </del>	loop	loop		-	
AA		-		353	ļ	<del> </del>	<del> </del>		<del> </del>	ļ	
SS		ļ		H8	<u> </u>	+-		435	435	<del> </del>	
AA		-			<u> </u>		-	H11	H11	<del> </del>	+-
SS		ļ			-	+-	442	442	HII	<del> </del>	+
AA		<del> </del>			<del></del>		H11	H11	<del> </del>	<b>-</b>	
SS				<u> </u>	<del> </del>		HII	455	<del> </del>	<del> </del>	+-
AA		+		<del> </del>	+		<del> </del>	H12	+	<del> </del>	+
SS			1				_l	I TIZ		ــــــــــــــــــــــــــــــــــــــ	

AA = Amino Acid

SS = Secondary Structure

Table 19
Coordination Structure of TR-β and GC

	(	Coord	linatio	on Struc			β and GC				
Coordina- tion Structure	Rı	R <sub>2</sub>	R <sub>3</sub>	R <sub>5</sub>	R <sub>6</sub>	R2	R3	R4	R5	R6	Х
Structure	-O-CH <sub>2</sub> CO <sub>2</sub> H	Н	CH <sub>3</sub>	CH <sub>3</sub>	Н	Н	CH(CH <sub>3</sub> )	OH	H	Н	CH <sub>2</sub>
ĀĀ	0 011200211		0-13	0.1.5			269				
SS		<del></del>					H3				
AA A	<del> </del>		272				1				
SS			H3								
AA			273	<del></del>	<del> </del>		273				
SS		<del> </del>	H3				H3	<b> </b>			
AA	<del> </del>		275					h			
SS	<del> </del>	<del> </del>	H3			_					
AA	<del>                                     </del>	<del> </del>	276	<del>                                     </del>			<del> </del>	276	276	276	_
SS	<del> </del>		H3	<del> </del>				H3	Н3	H3	
AA	279	279	1.2	<del>                                     </del>	<del> </del>		<del> </del>				
SS	H3	H3	<del>                                     </del>	<del> </del>	<del> </del>	<del>                                     </del>	<del> </del>	<del>                                     </del>	-		
AA	282	+	<del> </del>		1		<del> </del>	<del>                                     </del>		l	<b>†</b>
SS	H3	1		<del> </del>	<del> </del>		<del>                                     </del>				<u> </u>
AA	113	<del> </del>		310	<del> </del>	-	<del>                                     </del>	<del> </del>	310	310	
SS				H5-H6		<u> </u>			H5- H6	H5-H6	
AA	313	<del>                                     </del>		<del> </del>	313	<del> </del>					
SS	H5-H6	<del> </del>	<del>                                     </del>	<u> </u>	H5-H6		·				
AA	1		+		1		314	<u> </u>			
SS	<del></del>	1	+			<del> </del>	H5-H6	<del>                                     </del>			1
ĀĀ	316	1		<del>                                     </del>		<del>                                     </del>				í	
SS	Н5-Н6	1	+		<u> </u>	1	1	1			
AA	113 110	+	+	<del> </del>	<del> </del>	+	317				<u> </u>
SS	<del></del>	1	+	<del> </del>	<del></del>	$\vdash$	H5-H6				1
AA	320		<del>                                     </del>	<del> </del>		1					
SS	H5-H6	+	+	<del> </del>	<del>                                     </del>	+-	<del></del>				1
AA	329	<del>                                     </del>	+		1	+-	1	1	1		$\top$
SS	S3	+-	<del> </del>	<del>                                     </del>	+	╅			· ·		
AA	330	+	<del> </del>	330	<del>                                     </del>	<del>                                     </del>					1
SS	S3		<del> </del>	S3		1				i	
AA	331		1	<del>                                     </del>	<del>†                                    </del>	$\top$	+		1		
SS	loop	<del> </del>	1	+		1	<del> </del>	1			1
AA		+	+		-	+	344		1		1
SS			1	<del> </del>	<del>                                     </del>	1	loop	1	1		
AA		1	+	<del> </del>	1	1-	346	346		1	1
SS	<del> </del>		+		<del> </del>	<del> </del>	loop	loop		1	
AA	<del> </del>	1	<del> </del>	353	<del> </del>	+	1	· · · · ·	1		$\top$
SS	+	1	+	H8	1	+-		1	<b>†</b>	1	+
AA	-	+	~	1		+		435	435	<del>                                     </del>	<del>                                     </del>
125		+	+	+		+-	<del></del>	HII	HII	<del>                                     </del>	+
<del></del>	-	+	+	+		+-	<del></del>	455	<u>-</u>	+	1
SS		+	+	<del> </del>		+	1	H12	<del>                                     </del>	+	<del>                                     </del>

AA = Amino Acid

SS = Secondary Structure

WO 99/26966

PCT/US98/25296

#### APPENDIX 3

#### TR DMT.PDB

REMARK TR dmt full length numbering

REMARK

REMARK Rfactor 0.205 Rfree 0.227 5

REMARK Resolution 15. 2.2 all reflections

REMARK

REMARK Three cacodylate-modified cysteines (CYA)

REMARK Cya334, Cya380, Cya392

REMARK cacodylate modeled as single arsenic atom 10

REMARK

REMARK side chain of certain residues modeled as ALA due to poor density;

REMARK however, residue name reflects true residue for clarity

REMARK

REMARK clone obtained from Murray et. al. 15

REMARK deposited sequence confirmed,

REMARK differing from that reported by Thompson et. al.

REMARK in the following codons:

REMARK 281 Thr - Ala

20 REMARK 285 Lys - Glu

REMARK identical to that reported by Mitsuhashi et. al.

REMARK gb:RNTRAVI X07409

AUTH **JRNL** 

M.B. MURRAY, N.D.ZILZ,

N.L.MCCREARY, M.J.MACDONALD

25 **JRNL AUTH 2 H.C.TOWLE** 

TITL ISOLATION AND CHARACTERIZATION OF RAT CDNA **JRNL CLONES FOR TWO** 

TITL 2 DISTINCT THYROID HORMONE RECPTORS JRNL

JRNL REF JBC

45

V. 263 25 1988

AUTH C.C.THOMPSON, C.WEINBERGER, R.LEBO, R.M.EVANS 30 JRNL TITL IDENTIFICATION OF A NOVEL THYROID HORMONE JRNL RECEPTOR EXPRESSED

TITL 2 IN THE MAMMALIAN CENTRAL NERVOUS SYSTEM JRNL

V. 237 JRNL REF SCIENCE

1987 AUTH T.MITSUHASHI,G.TENNYSON,V.NIKODEM

35 JRNL TITL NUCLEOTIDE SEQUENCE OF NOVEL CDNAS GENERATED JRNL BY ALTERNATIVE

TITL 2 SPLICING OF A RAT THYROID HORMONE RECEPTOR JRNL GENE TRANSCRIPT

REF NUC. ACIDS. RES. V. 16 12 1988 40 JRNL

68.504 8.445 5.651 1.00 68.93 1 N ARG 157 **ATOM** 

67.886 9.543 6.398 1.00 56.98 ATOM 2 CA ARG 157

68.769 10.789 6.324 1.00 59.25 3 CB ARG 157 ATOM

70.147 10.632 6.932 1.00 58.90 ATOM 4 CG ARG 157

70.068 10.422 8.425 1.00 59.37 ATOM 5 CD ARG 157

71.392 10.446 9.036 1.00 63.94 6 NE ARG 157 ATOM

71.613 10.329 10.341 1.00 64.39 ATOM 7 CZ ARG 157

70.596 10.182 11.179 1.00 62.14 8 NH1 ARG 157 ATOM

```
72.855 10.365 10.808 1.00 65.56
             9 NH2 ARG 157
    ATOM
    ATOM
            10 C ARG 157
                             66.500 9.881 5.854 1.00 48.97
            11 O ARG 157
                             66.351 10.203 4.674 1.00 48.61
    ATOM
            12 N PRO 158
                             65.469 9.818 6.712 1.00 41.90
    ATOM
                             65.550 9.366 8.112 1.00 41.06
            13 CD PRO 158
    ATOM
                              64.083 10.114 6.333 1.00 39.34
            14 CA PRO 158
    ATOM
                             63.286 9.704 7.576 1.00 37.89
            15 CB PRO 158
    ATOM
            16 CG PRO 158
                             64.260 9.883 8.693 1.00 42.40
    ATOM
            17 C PRO 158
                             63.814 11.573 5.930 1.00 37.10
    ATOM
            18 O PRO 158
                             64.189 12.517 6.636 1.00 33.31
10
    ATOM
                             63.171 11.733 4.778 1.00 30.56
    ATOM
            19 N GLU 159
                             62.821 13.038 4.231 1.00 24.26
    ATOM
            20 CA GLU 159
            21 CB GLU 159
                              62.553 12.904 2.727 1.00 19.19
    ATOM
            22 CG GLU 159
                              63.788 12.677 1.874 1.00 20.60
    ATOM
                              64.407 13.971 1.390 1.00 26.54
            23 CD GLU 159
    ATOM
15
                              63.649 14.929 1.115 1.00 30.85
            24 OE1 GLU 159
    ATOM
                               65.649 14.027 1.268 1.00 28.35
    ATOM
            25 OE2 GLU 159
                             61.549 13.520 4.909 1.00 23.26
            26 C GLU. 159
    ATOM
            27 O GLU 159
                              60.906 12.765 5.643 1.00 26.86
    ATOM
                             61.200 14.806 4.729 1.00 22.72
            28 N PRO 160
20
    ATOM
                             61.981 15.916 4.153 1.00 17.87
    ATOM
            29 CD PRO 160
                              59.969 15.292 5.359 1.00 19.90
    ATOM
            30 CA PRO 160
            31 CB PRO 160
                             60.004 16.799 5.070 1.00 14.42
    ATOM
            32 CG PRO 160
                             61.465 17.109 4.919 1.00 12.87
    ATOM
                             58.747 14.623 4.701 1.00 23.68
            33 C PRO 160
    ATOM
25
                            58.730 14.383 3.491 1.00 24.72
            34 O PRO 160
    ATOM
                             57.749 14.281 5.506 1.00 22.19
            35 N THR 161
    ATOM
                             56.542 13.660 4.985 1.00 19.50
            36 CA THR 161
    ATOM
                              55.691 13.031 6.125 1.00 21.50
            37 CB THR 161
    ATOM
                              55.163 14.062 6.972 1.00 20.33
            38 OG1 THR 161
30
    ATOM
                               56.537 12.078 6.959 1.00 19.48
            39 CG2 THR 161
    ATOM
                              55.744 14.765 4.298 1.00 22.86
    ATOM
            40 C THR 161
                              56.040 15.949 4.481 1.00 27.68
    ATOM
            41 O THR 161
            42 N PRO 162
                              54.720 14.403 3.504 1.00 20.36
    ATOM
                              54.280 13.050 3.113 1.00 16.55
            43 CD PRO 162
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    ATOM
                              53.924 15.435 2.830 1.00 21.97
            44 CA PRO 162
    ATOM
                              52.780 14.633 2.210 1.00 18.17
            45 CB PRO 162
    ATOM
            46 CG PRO 162
                             53.422 13.316 1.905 1.00 18.01
     ATOM
            47 C PRO 162
                            53.399 16.467 3.826 1.00 22.56
     ATOM
                            53.461 17.675 3.567 1.00 21.73
            48 O PRO 162
40
    ATOM
                              52.912 15.976 4.967 1.00 25.28
            49 N GLU 163
     ATOM
                             52.357 16.816 6.030 1.00 26.64
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             50 CA GLU 163
                              51.743 15.962 7.144 1.00 30.22
             51 CB GLU 163
     ATOM
                               50.514 15.131 6.748 1.00 44.99
             52 CG GLU 163
     ATOM
             53 CD GLU 163
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45
                               50.016 13.660 4.929 1.00 52.48
             54 OE1 GLU 163
     ATOM
                               51.895 13.309 6.015 1.00 44.23
            55 OE2 GLU 163
     ATOM
            56 C GLU 163
                              53.414 17.731 6.634 1.00 27.65
     ATOM
            57 O GLU 163
                              53.114 18.862 7.034 1.00 29.30
     ATOM
                             54.646 17.235 6.712 1.00 21.89
            58 N GLU 164
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     ATOM
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WO 99/26966

PCT/US98/25296

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55.741 18.015 7.265 1.00 18.29
             59 CA GLU 164
    ATOM
                               56.901 17.109 7.657 1.00 14.78
    ATOM
             60 CB GLU 164
                               56.552 16.196 8.825 1.00 21.11
    ATOM
             61 CG GLU 164
             62 CD GLU 164
                               57.669 15.249 9.198 1.00 20.35
    ATOM
                               58.605 15.071 8.392 1.00 28.55
             63 OE1 GLU 164
    ATOM
                               57.610 14.677 10.302 1.00 28.25
             64 OE2 GLU 164
    ATOM
                              56.200 19.097 6.306 1.00 24.62
             65 C GLU 164
    ATOM
                              56.574 20.183 6.741 1.00 32.05
             66 O GLU 164
    ATOM
                              56.174 18.817 5.003 1.00 28.22
             67 N TRP 165
    ATOM
                               56.576 19.825 4.021 1.00 22.99
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             68 CA TRP 165
                               56.575 19.262 2.605 1.00 17.37
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             69 CB TRP 165
                               57.876 18.633 2.210 1.00 10.74
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             70 CG TRP 165
                               59.153 19.283 2.109 1.00 11.74
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             71 CD2 TRP 165
             72 CE2 TRP 165
                               60.075 18.319 1.648 1.00 9.97
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                               59.606 20.583 2.365 1.00 13.88
             73 CE3 TRP 165
    ATOM
15
                               58.074 17.343 1.832 1.00 9.17
             74 CD1 TRP 165
     ATOM
                               59.390 17.145 1.486 1.00 16.55
     ATOM
             75 NE1 TRP 165
                               61.427 18.613 1.436 1.00 13.37
             76 CZ2 TRP 165
     ATOM
                               60.954 20.874 2.156 1.00 16.15
             77 CZ3 TRP 165
     ATOM
                               61.846 19.892 1.696 1.00 17.42
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     ATOM
             78 CH2 TRP 165
                              55.634 21.015 4.115 1.00 21.44
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             79 C TRP 165
                              56.041 22.149 3.865 1.00 22.12
     ATOM
             80 O TRP 165
     ATOM
             81 N ASP 166
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                              53.369 21.796 4.621 1.00 25.77
             82 CA ASP 166
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                               51.972 21.196 4.808 1.00 26.02
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             84 CG ASP 166
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             85 OD1 ASP 166
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                                50.537 19.692 3.649 1.00 34.47
             86 OD2 ASP 166
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             87 C ASP 166
                              53.744 23.865 5.767 1.00 31.28
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             88 O ASP 166
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                              54.046 21.966 6.951 1.00 25.57
             89 N LEU 167
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                               54.439 22.640 8.187 1.00 28.28
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             90 CA LEU 167
                               54.854 21.624 9.256 1.00 32.80
     ATOM
             91 CB LEU 167
                               53.945 21.347 10.455 1.00 41.75
             92 CG LEU 167
     ATOM
                                54.765 20.640 11.532 1.00 39.15
     ATOM
             93 CD1 LEU 167
35
                                53.374 22.647 11.008 1.00 39.20
             94 CD2 LEU 167
     ATOM
                               55.636 23.532 7.902 1.00 22.19
             95 C LEU 167
     ATOM
                               55.671 24.700 8.302 1.00 29.51
             96 O LEU 167
     ATOM
                              56.610 22.957 7.206 1.00 15.01
             97 N ILE 168
     ATOM
             98 CA ILE 168
                               57.846 23.632 6.833 1.00 18.03
     ATOM
40
             99 CB ILE 168
                               58.756 22.668 6.040 1.00 11.37
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                                59.890 23.413 5.367 1.00 16.36
     ATOM
             100 CG2 ILE 168
                                59.289 21.580 6.975 1.00 21.63
     ATOM
             101 CG1 ILE 168
                                60.095 20.501 6.287 1.00 21.03
             102 CD1 ILE 168
     ATOM
                              57.579 24.897 6.022 1.00 22.54
             103 C ILE 168
45
     ATOM
                               58.155 25.948 6.300 1.00 24.88
     ATOM
             104 O ILE 168
                               56.682 24.800 5.045 1.00 25.70
             105 N HIS 169
     ATOM
                               56.337 25.934 4.190 1.00 21.28
             106 CA HIS 169
     ATOM
                               55.411 25.493 3.057 1.00 22.29
             107 CB HIS 169
     ATOM
                               56.047 24.543 2.091 1.00 23.11
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             108 CG HIS 169
     ATOM
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PCT/US98/25296

WO 99/26966

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ATOM

57.348 24.265 1.839 1.00 16.86 ATOM 109 CD2 HIS 169 55.312 23.721 1.263 1.00 25.30 110 ND1 HIS 169 56.130 22.974 0.546 1.00 15.89 ATOM 111 CEI HIS 169 57.371 23.283 0.878 1.00 25.38 112 NE2 HIS 169 ATOM 113 C HIS 169 55.664 27.048 4.976 1.00 18.32 ATOM 56.033 28.215 4.842 1.00 21.53 114 O HIS 169 ATOM 54.679 26.685 5.795 1.00 17.13 115 N VAL 170 ATOM 53.957 27.661 6.607 1.00 21.29 116 CA VAL 170 ATOM 52.808 26.991 7.399 1.00 24.33 117 CB VAL 170 ATOM 118 CG1 VAL 170 52.164 27.985 8.354 1.00 23.78 10 ATOM 51.760 26.439 6.435 1.00 18.87 ATOM 119 CG2 VAL 170 54.910 28.382 7.567 1.00 24.69 ATOM 120 C VAL 170 ATOM 121 O VAL 170 54.912 29.616 7.637 1.00 28.77 55.759 27.609 8.245 1.00 20.35 122 N ALA 171 ATOM 123 CA ALA 171 56.722 28.148 9.202 1.00 19.61 ATOM 15 57.393 27.013 9.977 1.00 17.52 124 CB ALA 171 ATOM 57.775 29.026 8.531 1.00 20.91 125 C ALA 171 ATOM 58.102 30.105 9.041 1.00 21.98 126 O ALA 171 ATOM 127 N THR 172 58.308 28.571 7.398 1.00 18.94 ATOM 59.313 29.342 6.668 1.00 19.55 128 CA THR 172 20 ATOM 59.820 28.594 5.413 1.00 20.49 ATOM 129 CB THR 172 60.394 27.336 5.795 1.00 20.66 ATOM 130 OG1 THR 172 60.894 29.418 4.702 1.00 20.44 ATOM 131 CG2 THR 172 58.730 30.697 6.254 1.00 23.26 ATOM 132 C THR 172 133 O THR 172 ATOM 59.403 31.724 6.334 1.00 24.32 25 134 N GLU 173 57.468 30.694 5.836 1.00 27.42 ATOM 56.797 31.922 5.434 1.00 27.68 135 CA GLU 173 ATOM 55.477 31.605 4.728 1.00 24.51 136 CB GLU 173 ATOM 137 CG GLU 173 54.652 32.836 4.338 1.00 39.69 ATOM 138 CD GLU 173 55.396 33.814 3.426 1.00 47.72 ATOM 30 55.019 35.009 3.417 1.00 48.26 ATOM 139 OE1 GLU 173 140 OE2 GLU 173 56.344 33.398 2.717 1.00 49.61 **ATOM** 56.557 32.834 6.641 1.00 25.68 ATOM 141 C GLU 173 ATOM 142 O GLU 173 56.773 34.046 6.559 1.00 23.39 56.119 32.245 7.755 1.00 25.19 143 N ALA 174 ATOM 35 144 CA ALA 174 55,863 32,989 8,993 1.00 22.25 **ATOM** 55.450 32.030 10.111 1.00 15.95 145 CB ALA 174 ATOM 57.125 33.747 9.391 1.00 23.22 ATOM 146 C ALA 174 57.076 34.918 9.768 1.00 24.52 ATOM 147 O ALA 174 148 N HIS 175 58.261 33.073 9.275 1.00 20.97 40 **ATOM** 149 CA HIS 175 59.544 33.665 9.606 1.00 19.55 ATOM 60.625 32.577 9.649 1.00 16.19 150 CB HIS 175 ATOM -62.016 33.104 9.835 1.00 18.89 ATOM 151 CG HIS 175 63.148 32.901 9.119 1.00 16.05 ATOM 152 CD2 HIS 175 153 ND1 HIS 175 62.359 33.962 10.859 1.00 13.83 ATOM 45 154 CE1 HIS 175 63.642 34.265 10.765 1.00 15.87 ATOM 64.143 33.635 9.718 1.00 19.19 155 NE2 HIS 175 **ATOM** 59.934 34.757 8.617 1.00 21.28 156 C HIS 175 ATOM 157 O HIS 175 60.274 35.869 9.014 1.00 25.12 ATOM 59.891 34.436 7.329 1.00 26.73 158 N ARG 176

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60.266 35.387 6.292 1.00 27.13
    ATOM
            159 CA ARG 176
                                60.156 34.748 4.914 1.00 36.00
            160 CB ARG 176
    ATOM
            161 CG ARG 176
                                61.286 33.795 4.602 1.00 43.20
    ATOM
            162 CD ARG 176
                                61.197 33.334 3.170 1.00 50.07
    ATOM
            163 NE ARG 176
                                62.316 32.477 2.813 1.00 58.20
    ATOM
                                62.266 31.548 1.867 1.00 67.22
            164 CZ ARG 176
    ATOM
                                61.143 31.358 1.182 1.00 67.62
    ATOM
            165 NH1 ARG 176
                                63.336 30.806 1.612 1.00 70.56
            166 NH2 ARG 176
    ATOM
                               59.487 36.688 6.325 1.00 23.97
            167 C ARG 176
    ATOM
                               60.073 37.760 6.209 1.00 24.52
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    ATOM
            168 O ARG 176
    ATOM
            169 N SER 177
                              58.177 36.598 6.515 1.00 23.60
            170 CA SER 177
                               57.341 37.789 6.565 1.00 26.36
    ATOM
            171 CB SER 177
                               55.865 37.407 6.439 1.00 21.93
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                              55.495 36.459 7.423 1.00 25.97 57.557 38.623 7.829 1.00 28.76
            172 OG SER 177
    ATOM
            173 C SER 177
    ATOM
15
                              57.084 39.761 7.907 1.00 33.09
            174 O SER 177
    ATOM
                               58.257 38.062 8.815 1.00 25.52
            175 N THR 178
    ATOM
                                58.508 38.772 10.064 1.00 18.93
            176 CA THR 178
    ATOM
                                57.828 38.064 11.258 1.00 21.81
    ATOM
            177 CB THR 178
    ATOM
            178 OG1 THR 178
                                58.348 36.736 11.394 1.00 24.18
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            179 CG2 THR 178
                                56.330 37.971 11.032 1.00 13.81
    ATOM
                               59.993 38.967 10.358 1.00 20.69
            180 C THR 178
    ATOM
            181 O THR 178
                               60.373 39.407 11.448 1.00 20.56
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            182 N ASN 179
                               60.837 38.645 9.385 1.00 23.68
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                                62.275 38.802 9.555 1.00 28.22
            183 CA ASN 179
25
    ATOM
                                63.022 37.627 8.927 1.00 27.45
            184 CB ASN 179
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                                64.460 37.529 9.402 1.00 33.98
            185 CG ASN 179
    ATOM
                                65.342 37.131 8.644 1.00 42.72
            186 OD1 ASN 179
     ATOM
                                64.702 37.865 10.667 1.00 31.14
            187 ND2 ASN 179
     ATOM
            188 C ASN 179
                               62.689 40.115 8.902 1.00 34.47
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    ATOM
                               62.832 40.200 7.678 1.00 36.54
     ATOM
            189 O ASN 179
                               62.874 41.135 9.735 1.00 37.39
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            190 N ALA 180
            191 CA ALA 180
                                63.235 42.479 9.292 1.00 33.71
     ATOM
                                63.555 43.352 10.494 1.00 31.57
            192 CB ALA 180
     ATOM
                               64.375 42.545 8.284 1.00 37.87
            193 C ALA 180
     ATOM
35
                               65.458 42.018 8.525 1.00 35.26
            194 O ALA 180
     ATOM
                               64.095 43.187 7.150 1.00 40.55
     ATOM
            195 N GLN 181
                              65.049 43.391 6.057 1.00 42.95
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            196 CA GLN 181
            197 CB GLN 181
                                66.344 44.043 6.570 1.00 45.47
     ATOM
            198 CG GLN 181
                                66.144 45.326 7.383 1.00 52.70
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     ATOM
                                65.351 46.399 6.650 1.00 55.03
            199 CD GLN 181
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                                65.270 46.412 5.421 1.00 59.56
     ATOM 200 OE1 GLN 181
            201 NE2 GLN 181
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                               65.391 42.176 5.197 1.00 44.27
            202 C GLN 181
     ATOM
            203 O GLN 181
                               66.181 42.291 4.251 1.00 46.47
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                                65.054 39.815 4.742 1.00 42.63
            205 CA GLY 182
     ATOM
            206 C GLY 182
207 O GLY 182
                               66.522 39.584 4.427 1.00 47.40
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                               67.382 39.691 5.306 1.00 49.38
     ATOM
                               66.816 39.297 3.163 1.00 49.46
            208 N SER 183
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68.189 39.061 2.733 1.00 54.13
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                               68.208 38.225 1.449 1.00 55.08
           210 CB SER 183
    ATOM
            211 OG SER 183
                               67.197 38.647 0.546 1.00 63.54
            212 C SER 183
                              68.949 40.369 2.532 1.00 54.84
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           213 O SER 183
                              70.175 40.373 2.407 1.00 56.90
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                             68.223 41.482 2.535 1.00 55.77
           214 N HIS 184
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                             68.854 42.775 2.342 1.00 57.78
            215 CA HIS 184
    ATOM
                             69.605 43.296 3.556 1.00 59.09
    ATOM
            216 C HIS 184
                              70.312 44.301 3.454 1.00 60.34
            217 O HIS 184
    ATOM
                              69.502 42.597 4.686 1.00 55.60
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    ATOM
            218 N TRP 185
                              70.159 43.020 5.923 1.00 53.73
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            219 CA TRP 185
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            220 CB TRP 185
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            221 CG TRP 185
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                               72.091 40.419 7.269 1.00 47.38
            222 CD2 TRP 185
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                               72.390 39.094 6.888 1.00 40.29
            223 CE2 TRP 185
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15
                               73.071 41.169 7.937 1.00 45.43
            224 CE3 TRP 185
    ATOM
                               70.301 39.554 6.234 1.00 49.87
            225 CD1 TRP 185
    ATOM
                               71.280 38.589 6.262 1.00 48.02
            226 NEI TRP 185
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                               73.628 38.496 7.154 1.00 38.65
            227 CZ2 TRP 185
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                               74.304 40.573 8.201 1.00 43.26
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    ATOM
           228 CZ3 TRP 185
                               74.570 39.250 7.807 1.00 40.00
    ATOM 229 CH2 TRP 185
                              71.638 43.386 5.800 1.00 55.99
    ATOM 230 C TRP 185
                              72.089 44.359 6.401 1.00 52.84
    ATOM 231 O TRP 185
            232 N LYS 186
                              72.389 42.614 5.021 1.00 59.15
    ATOM
                               73.818 42.863 4.843 1.00 64.01
            233 CA LYS 186
    ATOM
25
                               74.466 41.688 4.091 1.00 64.67
            234 CB LYS 186
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                               75.943 41.868 3.729 1.00 65.58
            235 CG LYS 186
    ATOM
                               76.817 42.181 4.946 1.00 62.03
            236 CD LYS 186
     ATOM
                               78.238 42.512 4.515 1.00 61.52
           237 CE LYS 186
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                               78.988 43.243 5.579 1.00 61.67
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            238 NZ LYS 186
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           239 C LYS 186
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                              75.164 44.816 4.432 1.00 68.66
           240 O LYS 186
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                               73.221 44.678 3.316 1.00 68.99
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            241 N GLN 187
                               73.431 45.939 2.612 1.00 69.65
     ATOM 242 CA GLN 187
     ATOM 243 CB GLN 187
                               72.880 45.867 1.180 1.00 73.76
35
                               73.632 44.935 0.237 1.00 78.61
            244 CG GLN 187
     ATOM
                               73.368 43.471 0.525 1.00 84.96
            245 CD GLN 187
     ATOM
                                74.203 42.782 1.109 1.00 87.73
            246 OE1 GLN 187
     ATOM
                                72.197 42.989 0.122 1.00 84.98
           247 NE2 GLN 187
     ATOM
                               72.817 47.141 3.323 1.00 69.16
     ATOM
           248 C GLN 187
40
                               73.379 48.235 3.299 1.00 71.39
           249 O GLN 187
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                               71.666 46.936 3.953 1.00 65.82
     ATOM 250 N ARG 188
                               70.961 48.014 4.639 1.00 65.00
     ATOM 251 CA ARG 188
                               69.458 47.739 4.591 1.00 66.20
            252 CB ARG 188
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                                68.957 47.483 3.181 1.00 70.30
     ATOM 253 CG ARG 188
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                                67.463 47.212 3.132 1.00 78.59
     ATOM 254 CD ARG 188
                                67.003 47.008 1.760 1.00 87.71
            255 NE ARG 188
     ATOM
                               67.011 47.946 0.814 1.00 94.10
     ATOM 256 CZ ARG 188
                               67.453 49.171 1.081 1.00 97.26
            257 NH1 ARG 188
     ATOM
                                66.589 47.657 -0.409 1.00 94.07
     ATOM 258 NH2 ARG 188
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71.409 48.286 6.077 1.00 65.39
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    ATOM 260 O ARG 188
                              72.372 47.506 6.561 1.00 64.28
            261 N ARG 189
    ATOM
                               72.882 47.654 7.922 1.00 60.75
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    ATOM
            263 CB ARG 189
                               73.691 46.409 8.321 1.00 56.87
    ATOM
                               75.050 46.308 7.630 1.00 59.52
            264 CG ARG 189
    ATOM
                               75.580 44.891 7.589 1.00 55.86
            265 CD ARG 189
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                               75.874 44.348 8.907 1.00 55.48
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                               77.055 43.849 9.257 1.00 61.38
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    ATOM
                                78.057 43.832 8.388 1.00 62.54
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    ATOM
           268 NH1 ARG 189
           269 NH2 ARG 189
                                77.225 43.328 10.465 1.00 62.20
    ATOM
    ATOM 270 C ARG 189
                              73.747 48.907 8.082 1.00 60.91
    ATOM 271 O ARG 189
                              74.548 49.245 7.207 1.00 60.67
           272 N LYS 190
                              73.575 49.591 9.207 1.00 59.06
    ATOM
                               74.340 50.790 9.521 1.00 55.00
           273 CA LYS 190
    ATOM
15
                              73.423 52.008 9.582 1.00 55.45
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                              74.991 50.542 10.875 1.00 51.52
            275 C LYS 190
    ATOM
            276 O LYS 190
                            74.320 50.144 11.830 1.00 51.68
    ATOM
            277 N PHE 191
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    ATOM
                               77.037 50.508 12.186 1.00 50.17
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           278 CA PHE 191
20
                               78.546 50.571 11.943 1.00 48.38
            279 CB PHE 191
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                               79.090 49.423 11.142 1.00 49.66
            280 CG PHE 191
    ATOM
                               78.873 49.348 9.768 1.00 51.03
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    ATOM
            282 CD2 PHE 191
                               79.845 48.429 11.759 1.00 46.28
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           283 CE1 PHE 191
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    ATOM
                               80.379 47.377 11.021 1.00 47.26
            284 CE2 PHE 191
    ATOM
                               80.158 47.311 9.646 1.00 48.48
            285 CZ PHE 191
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                              76.663 51.534 13.248 1.00 48.61
            286 C PHE 191
    ATOM
            287 O PHE 191
                              76.507 52.720 12.952 1.00 50.38
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                              76.488 51.068 14.479 1.00 47.31
           288 N LEU 192
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           289 CA LEU 192
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    ATOM 291 CG LEU 192
                                74.048 52.590 17.786 1.00 28.37
     ATOM 292 CD1 LEU 192
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     ATOM 293 CD2 LEU 192
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                              77.447 52.760 15.800 1.00 42.28
     ATOM 294 C LEU 192
                              78.528 52.179 15.932 1.00 39.71
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           296 N PRO 193
     ATOM
           297 CD PRO 193
                               76.095 54.865 15.617 1.00 43.82
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     ATOM 298 CA PRO 193
                               78.493 55.006 15.973 1.00 43.14
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                               77.820 56.306 16.400 1.00 44.37
     ATOM 299 CB PRO 193
                               76.571 56.308 15.565 1.00 41.66
     ATOM 300 CG PRO 193
     ATOM 301 C PRO 193
                              79.476 54.498 17.028 1.00 43.34
            302 O PRO 193
                               79.103 54.296 18.182 1.00 45.18
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     ATOM
                              81.781 53.804 17.512 1.00 47.20
            304 CA ASP 194
     ATOM
                               83.108 53.732 16.761 1.00 41.89
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                              81.962 54.511 18.866 1.00 51.99
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     ATOM
                              82.636 53.986 19.752 1.00 54.04
            307 O ASP 194
     ATOM
                              81.381 55.698 19.025 1.00 55.21
            308 N ASP 195
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PCT/US98/25296

## WO 99/26966

	ATOM	309 CA ASP 195	81.489 56.428 20.288 1.00 57.50
	ATOM	310 CB ASP 195	81.423 57.948 20.061 1.00 60.04
	ATOM	311 CG ASP 195	80.123 58.398 19.406 1.00 68.39
	ATOM	312 OD1 ASP 195	79.211 58.847 20.136 1.00 69.46
5	ATOM	313 OD2 ASP 195	80.020 58.322 18.162 1.00 72.91
	ATOM	314 C ASP 195	80.410 55.976 21.280 1.00 58.05
	ATOM	315 O ASP 195	80.540 56.180 22.491 1.00 58.97
	ATOM	316 N ILE 196	79.349 55.363 20.759 1.00 56.06
	<b>ATOM</b>	317 CA ILE 196	78.247 54.863 21.580 1.00 50.48
10	<b>ATOM</b>	318 CB ILE 196	76.930 54.762 20.766 1.00 45.82
	ATOM	319 CG2 ILE 196	75.818 54.166 21.621 1.00 44.04
	<b>ATOM</b>	320 CG1 ILE 196	76.517 56.147 20.261 1.00 44.27
	<b>ATOM</b>	321 CD1 ILE 196	75.179 56.171 19.541 1.00 45.25
	ATOM	322 C ILE 196	78.603 53.484 22.135 1.00 47.66
15	ATOM	323 O ILE 196	79.138 52.636 21.419 1.00 43.96
	ATOM	324 N GLY 197	78.309 53.269 23.414 1.00 46.29
	ATOM	325 CA GLY 197	78.608 51.995 24.045 1.00 48.03
	ATOM	326 C GLY 197	79.978 51.963 24.692 1.00 50.42
	ATOM	327 O GLY 197	80.463 50.902 25.070 1.00 46.66
20	ATOM	328 N GLN 198	80.583 53.137 24.854 1.00 56.94
	<b>ATOM</b>	329 CA GLN 198	81.910 53.259 25.454 1.00 59.51
	ATOM	330 CB GLN 198	82.751 54.257 24.649 1.00 62.53
	ATOM	331 CG GLN 198	83.232 53.718 23.316 1.00 69.39
	<b>ATOM</b>	332 CD GLN 198	84.088 52.484 23.483 1.00 76.76
25	ATOM	333 OE1 GLN 198	83.745 51.399 22.996 1.00 81.73
	<b>ATOM</b>	334 NE2 GLN 198	85.205 52.632 24.192 1.00 78.09
	ATOM	335 C GLN 198	81.915 53.678 26.922 1.00 57.56
	ATOM	336 O GLN 198	82.946 53.584 27.588 1.00 57.71
	<b>ATOM</b>	337 N SER 199	80.770 54.128 27.425 1.00 54.11
30	ATOM	338 CA SER 199	80.676 54.600 28.800 1.00 46.28
	ATOM	339 CB SER 199	80.243 56.067 28.777 1.00 50.28
	ATOM	340 OG SER 199	80.935 56.776 27.757 1.00 50.95
	ATOM	341 C SER 199	79.776 53.805 29.757 1.00 40.19
	ATOM	342 O SER 199	78.680 54.252 30.102 1.00 39.26
35	<b>ATOM</b>	343 N PRO 200	80.236 52.629 30.214 1.00 35.63
	ATOM	344 CD PRO 200	81.530 52.011 29.904 1.00 34.88
	ATOM	345 CA PRO 200	79.464 51.789 31.139 1.00 37.54
	ATOM	346 CB PRO 200	80.223 50.457 31.124 1.00 29.86
	ATOM	347 CG PRO 200	81.207 50.570 29.995 1.00 34.29
40	ATOM	348 C PRO 200	79.521 52.416 32.532 1.00 44.63
	ATOM	349 O PRO 200	80.443 52.137 33.300 1.00 47.80
	ATOM	350 N ILE 201	78.532 53.241 32.867 1.00 49.57
	ATOM	351 CA ILE 201	78.525 53.924 34.158 1.00 49.15
	ATOM	352 CB ILE 201	78.213 55.426 33.990 1.00 49.19
45	<b>ATOM</b>	353 CG2 ILE 201	78.429 56.150 35.306 1.00 53.37
	<b>ATOM</b>	354 CG1 ILE 201	79.137 56.037 32.934 1.00 52.55
	<b>ATOM</b>	355 CD1 ILE 201	78.811 57.471 32.586 1.00 55.26
	<b>ATOM</b>	356 C ILE 201	77.625 53.352 35.254 1.00 49.88
	ATOM	357 O ILE 201	78.044 53.250 36.408 1.00 50.20
50	ATOM	358 N VAL 202	76.384 53.014 34.920 1.00 47.85

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            360 CB VAL 202
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            361 CG1 VAL 202
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            362 CG2 VAL 202
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                               75.954 51.093 36.373 1.00 50.57
            363 C VAL 202
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                               76.296 50.249 35.545 1.00 49.50
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            365 N SER 203
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                               76.490 49.609 38.223 1.00 59.26
            366 CA SER 203
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                               77.067 49.809 39.628 1.00 64.88
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                               76.127 50.428 40.492 1.00 75.47
            368 OG SER 203
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            369 C SER 203
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            370 O SER 203
                               75.923 47.283 37.958 1.00 52.29
            371 N MET 204
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            372 CA MET 204
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            373 CB MET 204
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            374 CG MET 204
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            375 SD MET 204
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                                71.947 44.785 35.241 1.00 39.19
            376 CE MET 204
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            377 C MET 204
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            378 O MET 204
                               76.892 45.020 39.062 1.00 52.25
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                               74.816 44.329 39.605 1.00 47.73
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                                73.344 44.414 39.549 1.00 48.94
            380 CD PRO 205
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            381 CA PRO 205
                                75.250 43.326 40.580 1.00 47.34
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             382 CB PRO 205
                                73.982 42.513 40.810 1.00 49.44
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                                72.907 43.562 40.725 1.00 50.62
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25
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                               76.431 42.442 40.168 1.00 47.12
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                               77.299 42.160 40.990 1.00 51.21
             385 O PRO 205
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             386 N ASP 206
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30
                                76.598 40.967 36.107 1.00 57.34
             389 CG ASP 206
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                                77.056 42.095 35.811 1.00 52.21
     ATOM
             390 OD1 ASP 206
                                75.719 40.397 35.423 1.00 59.16
             391 OD2 ASP 206
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                               78.902 41.843 38.093 1.00 48.70
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             392 C ASP 206
                               79.862 41.171 37.715 1.00 49.75
             393 O ASP 206
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35
                               78.946 43.168 38.161 1.00 47.54
             394 N GLY 207
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                                80.174 43.869 37.820 1.00 49.23
             395 CA GLY 207
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             396 C GLY 207
                               80.169 44.585 36.482 1.00 51.96
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                               80.783 45.645 36.348 1.00 56.32
             397 O GLY 207
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             398 N ASP 208
                               79.510 44.005 35.481 1.00 52.50
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                                79.435 44.624 34.157 1.00 48.00
             399 CA ASP 208
     ATOM
                                78.968 43.609 33.115 1.00 53.23
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             400 CB ASP 208
             401 CG ASP 208
                                80.038 42.592 32.774 1.00 53.17
     ATOM
             402 OD1 ASP 208
                                81.130 43.006 32.335 1.00 57.42
     ATOM
                                79.787 41.380 32.942 1.00 55.64
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             403 OD2 ASP 208
45
                               78.497 45.823 34.187 1.00 46.68
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             404 C ASP 208
                               77.283 45.671 34.332 1.00 45.81
             405 O ASP 208
     ATOM
                               79.075 47.014 34.077 1.00 45.95
             406 N LYS 209
     ATOM
                                78.313 48.257 34.115 1.00 45.87
     ATOM
             407 CA LYS 209
                                79.235 49.418 34.478 1.00 46.90
             408 CB LYS 209
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	ATOM	409 C LYS 209	77.561 48.546 32.812 1.00 41.17
	ATOM	410 O LYS 209	77.951 48.074 31.745 1.00 39.51
	ATOM	411 N VAL 210	76.500 49.344 32.916 1.00 39.35
	ATOM	412 CA VAL 210	75.652 49.713 31.782 1.00 38.03
5	ATOM	413 CB VAL 210	74.136 49.584 32.140 1.00 32.13
_	ATOM	414 CG1 VAL 210	73.269 49.926 30.937 1.00 27.92
	ATOM	415 CG2 VAL 210	73.818 48.183 32.627 1.00 29.43
	ATOM	416 C VAL 210	75.895 51.134 31.263 1.00 38.68
	ATOM	417 O VAL 210	76.090 52.079 32.038 1.00 39.57
10	ATOM	418 N ASP 211	75.848 51.272 29.942 1.00 39.19
	<b>ATOM</b>	419 CA ASP 211	76.019 52.544 29.254 1.00 38.39
	<b>ATOM</b>	420 CB ASP 211	76.794 52.327 27.946 1.00 40.36
	<b>ATOM</b>	421 CG ASP 211	77.051 53.620 27.177 1.00 36.85
	<b>ATOM</b>	422 OD1 ASP 211	76.193 54.528 27.167 1.00 37.95
15	<b>ATOM</b>	423 OD2 ASP 211	78.121 53.716 26.553 1.00 33.87
	<b>ATOM</b>	424 C ASP 211	74.601 53.040 28.958 1.00 40.60
	ATOM ·		73.919 52.517 28.073 1.00 40.36
	ATOM	426 N LEU 212	74.185 54.074 29.680 1.00 41.55
	ATOM	427 CA LEU 212	72.854 54.664 29.552 1.00 38.39
20	ATOM	428 CB LEU 212	72.759 55.883 30.467 1.00 40.93
	ATOM	429 CG LEU 212	71.575 55.979 31.428 1.00 45.32
	ATOM	430 CD1 LEU 212	71.271 54.626 32.047 1.00 43.83
	ATOM	431 CD2 LEU 212	71.900 57.007 32.502 1.00 44.93
	ATOM	432 C LEU 212	72.448 55.050 28.133 1.00 37.61
25	ATOM	433 O LEU 212	71.318 54.805 27.719 1.00 33.71
	ATOM	434 N GLU 213	73.360 55.670 27.393 1.00 41.23 73.068 56.084 26.023 1.00 43.48
	ATOM	435 CA GLU 213	
	ATOM	436 CB GLU 213	74.181 56.986 25.481 1.00 47.66 73.919 57.494 24.065 1.00 56.87
20	ATOM	437 CG GLU 213	
30	ATOM	438 CD GLU 213 439 OE1 GLU 213	
	ATOM	439 OE1 GLU 213 440 OE2 GLU 213	76.258 57.996 23.924 1.00 60.37 74.921 58.894 22.423 1.00 61.13
	ATOM ATOM	441 C GLU 213	72.889 54.880 25.102 1.00 39.29
	ATOM	442 O GLU 213	71.965 54.841 24.290 1.00 36.66
35	ATOM	442 O GLO 213 443 N ALA 214	73,785 53.906 25.233 1.00 36.33
22	ATOM	444 CA ALA 214	73.739 52.693 24.422 1.00 34.89
	ATOM	445 CB ALA 214	74.946 51.817 24.711 1.00 30.70
	ATOM	446 C ALA 214	72.454 51.938 24.718 1.00 31.96
	ATOM	447 O ALA 214	71.739 51.523 23.804 1.00 33.93
40	ATOM	448 N PHE 215	72.151 51.798 26.003 1.00 28.47
	ATOM	449 CA PHE 215	70.947 51.116 26.445 1.00 29.74
	ATOM	450 CB PHE 215	70.819 51.223 27.962 1.00 23.73
	ATOM	451 CG PHE 215	69.589 50.568 28.515 1.00 22.71
	ATOM	452 CD1 PHE 215	69.603 49.220 28.858 1.00 22.53
45	<b>ATOM</b>	453 CD2 PHE 215	68.423 51.301 28.712 1.00 19.74
	ATOM	454 CE1 PHE 215	68.477 48.606 29.391 1.00 20.75
	ATOM	455 CE2 PHE 215	67.290 50.698 29.245 1.00 21.02
	ATOM	456 CZ PHE 215	67.318 49.346 29.586 1.00 19.50
	ATOM	457 C PHE 215	69.730 51.742 25.771 1.00 34.64
50	ATOM	458 O PHE 215	68.872 51.034 25.239 1.00 39.86

WO 99/26966

PCT/US98/25296

	ATOM	459 N SER 216	69.677 53.071 25.771 1.00 34.78
	ATOM	460 CA SER 216	68.572 53.801 25.160 1.00 36.01
	ATOM	461 CB SER 216	68.762 55.302 25.366 1.00 37.36
	ATOM	462 OG SER 216	67.537 55.987 25.193 1.00 48.33
5	ATOM	463 C SER 216	68.458 53.475 23.664 1.00 37.06
,	ATOM	464 O SER 216	67.358 53.250 23.148 1.00 33.23
	ATOM	465 N GLU 217	69.601 53.410 22.986 1.00 36.25
	ATOM	466 CA GLU 217	69.645 53.091 21.562 1.00 36.99
	ATOM	467 CB GLU 217	71.092 53.104 21.064 1.00 37.10
10	ATOM	468 CG GLU 217	71.682 54.491 20.912 1.00 44.30
10	ATOM	469 CD GLU 217	71.016 55.284 19.802 1.00 51.30
	ATOM	470 OE1 GLU 217	71.439 55.142 18.633 1.00 57.25
	ATOM	470 OE1 GLU 217	70.070 56.046 20.096 1.00 52.50
	ATOM	471 OE2 GLO 217 472 C GLU 217	69.019 51.726 21.286 1.00 36.93
15		472 C GLU 217 473 O GLU 217	68.191 51.577 20.381 1.00 41.06
15	ATOM	474 N PHE 218	69.395 50.740 22.093 1.00 30.27
	ATOM	474 N PHE 218	68.875 49.388 21.947 1.00 27.20
	ATOM	476 CB PHE 218	69.679 48.421 22.814 1.00 28.10
	ATOM	476 CB PHE 218	71.124 48.330 22.428 1.00 24.84
20	ATOM		72.117 48.286 23.398 1.00 21.78
20	ATOM	478 CD1 PHE 218 479 CD2 PHE 218	71.495 48.301 21.087 1.00 24.78
	ATOM		73.458 48.215 23.040 1.00 24.08
	ATOM ATOM		72.834 48.230 20.719 1.00 25.33
			73.818 48.187 21.697 1.00 25.04
25	ATOM ATOM	482 CZ PHE 218 483 C PHE 218	67.381 49.281 22.261 1.00 28.23
25			66,639 48.605 21.543 1.00 33.52
	ATOM		66.927 49.961 23.310 1.00 27.24
	ATOM		65.515 49.913 23.666 1.00 29.28
	ATOM	486 CA THR 219 487 CB THR 219	65,238 50.533 25.052 1.00 30.97
20	ATOM		65.724 51.880 25.090 1.00 35.50
30	ATOM		65.901 49.712 26.149 1.00 30.78
	ATOM	=	64.660 50.612 22.615 1.00 33.29
	ATOM		63.473 50.317 22.474 1.00 36.85
	ATOM		65.276 51.515 21.860 1.00 35.23
25	ATOM		64.579 52.253 20.816 1.00 38.97
35	ATOM		65.506 53.334 20.236 1.00 44.67
	ATOM		64.805 54.491 19.513 1.00 58.02
	ATOM	495 CG LYS 220	64.406 54.130 18.079 1.00 68.57
	ATOM	496 CD LYS 220 497 CE LYS 220	63.732 55.296 17.347 1.00 70.50
40	ATOM		
40	ATOM	498 NZ LYS 220	62.395 55.668 17.905 1.00 66.08 64.112 51.289 19.721 1.00 38.48
	ATOM	499 C LYS 220	63.021 51.446 19.173 1.00 37.18
	ATOM	500 O LYS 220	64.917 50.270 19.432 1.00 36.19
	ATOM	501 N ILE 221	64.563 49.305 18.394 1.00 36.77
4.5	ATOM	502 CA ILE 221	
45	ATOM	503 CB ILE 221	65.756 48.996 17.457 1.00 34.41
	ATOM	504 CG2 ILE 221	66.270 50.276 16.814 1.00 38.54
	ATOM	505 CG1 ILE 221	66.864 48.267 18.221 1.00 32.93
	ATOM	506 CD1 ILE 221	67.984 47.752 17.338 1.00 31.12 64.002 47.971 18.888 1.00 38.22
50	ATOM	507 C ILE 221	
50	ATOM	508 O ILE 221	63.499 47.181 18.089 1.00 38.90

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64.048 47.719 20.191 1.00 35.75
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                              63.557 46.446 20.702 1.00 31.77
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    ATOM
            511 CB ILE 222
                              64.086 46.152 22.130 1.00 33.14
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                               63.203 46.813 23.183 1.00 24.60
            512 CG2 ILE 222
    ATOM
                               64.147 44.638 22.350 1.00 32.60
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    ATOM
                               64.860 44.226 23.609 1.00 34.52
            514 CD1 ILE 222
    ATOM
                             62.042 46.240 20.624 1.00 32.56
           515 C ILE 222
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    ATOM 516 O ILE 222
                             61.581 45.109 20.452 1.00 35.74
                              61.262 47.313 20.720 1.00 29.43
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                               59,806 47.170 20.651 1.00 33.57
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                               59.075 48.514 20.903 1.00 38.99
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    ATOM
                               59.422 49.010 22.205 1.00 41.23
            520 OG1 THR 223
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            521 CG2 THR 223
                              59.355 46.528 19.325 1.00 31.45
            522 C THR 223
    ATOM
                               58.571 45.571 19.334 1.00 26.77
            523 O THR 223
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15
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            524 N PRO 224
                               60.570 48.306 17.950 1.00 30.11
            525 CD PRO 224
    ATOM
                               59.424 46.462 16.891 1.00 30.38
            526 CA PRO 224
    ATOM
            527 CB PRO 224
                               60.149 47.336 15.865 1.00 30.09
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                              60.200 48.659 16.530 1.00 31.86
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            529 C PRO 224
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                              59.147 44.153 16.295 1.00 32.52
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                               61.090 44.734 17.285 1.00 22.63
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                               60.752 42.416 18.026 1.00 23.53
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            536 N ILE 226
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                              58.112 41.768 19.251 1.00 21.28
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     ATOM 543 O ILE 226
                              57.553 40.670 19.256 1.00 23.75
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                               57.629 42.821 18.598 1.00 24.46
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            546 CB THR 227
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                                55.772 45.039 18.345 1.00 35.43
            547 OG1 THR 227
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                                54.776 44.049 16.388 1.00 29.01
            548 CG2 THR 227
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     ATOM
                               56.508 41.728 16.691 1.00 22.85
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            550 O THR 227
                               55,589 40,939 16,469 1,00 22,84
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                               57.647 41.713 16.004 1.00 16.09
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            554 CG ARG 228
                                60.309 42.447 12.422 1.00 20.90
            555 CD ARG 228
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           556 NE ARG 228
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                                61.729 44.328 13.973 1.00 36.35
     ATOM 558 NH1 ARG 228
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63.404 42.807 14.370 1.00 32.78
            559 NH2 ARG 228
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            560 C ARG 228
    ATOM
                               57.477 38.407 14.704 1.00 24.47
            561 O ARG 228
    ATOM
                               58.304 39.128 16.675 1.00 20.00
            562 N VAL 229
    ATOM
                               58.319 37.793 17.266 1.00 18.39
            563 CA VAL 229
    ATOM
                               59.103 37.745 18.606 1.00 19.20
           564 CB VAL 229
    ATOM
                                58.938 36.382 19.265 1.00 14.19
    ATOM 565 CG1 VAL 229
                                60.581 38.001 18.356 1.00 14.81
    ATOM 566 CG2 VAL 229
           567 C VAL 229
568 O VAL 229
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                               56.499 36.227 17.212 1.00 20.04
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            569 N VAL 230
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            570 CA VAL 230
    ATOM
                               53.930 39.185 18.912 1.00 22.15
    ATOM 571 CB VAL 230
                                52.452 38.862 19.113 1.00 15.66
            572 CG1 VAL 230
    ATOM
            573 CG2 VAL 230
                                54.592 39.522 20.248 1.00 21.05
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15
                               53.967 37.660 16.917 1.00 26.17
    ATOM 574 C VAL 230
                               53:188 36.704 16.836 1.00 28.01
            575 O VAL 230
    ATOM
                               54.288 38.426 15.873 1.00 25.07
           576 N ASP 231
     ATOM
                               53.714 38.216 14.542 1.00 26.10
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D	395 1.00 33.49
	376 1.00 40.13
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ATOM 617 O LYS 235 51.417 30.838 10.6	
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ATOM 619 CA LEU 236 51.722 50.665 12.5	
ATOM 620 CB LEU 236 52.944 29.080 14.	
ATOM 621 CG LEU 236 54.373 29.516 13	
ATOM 622 CD1 LEU 236 55.299 29.054 14	
15 ATOM 623 CD2 LEU 236 54.811 28.942 12 ATOM 624 C LEU 236 50.520 28.891 13.4	
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ATOM 625 0 LEG 230 49.930 29.333 14.3 ATOM 626 N PRO 237 50.012 27.895 12.3	
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ATOM 642 CA PHE 239 49.917 28.619 18	.117 1.00 41.11
	.788 1.00 34.80
ATOM 644 CG PHE 239 51.336 30.607 18	
ATOM 645 CD1 PHE 239 52.127 30.332 19	
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ATOM 647 CE1 PHE 239 52.368 31.307 20	0.896 1.00 30.28
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	.750 1.00 29.00
ATOM 649 CZ PHE 239 51.813 32.576 20	.750 1.00 29.00
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3	.750 1.00 29.00 337 1.00 35.65
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3 ATOM 651 O PHE 239 48.151 29.521 19.4 ATOM 652 N SER 240 48.133 30.037 17.3	.750 1.00 29.00 337 1.00 35.65 457 1.00 30.27 272 1.00 36.49
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3 ATOM 651 O PHE 239 48.151 29.521 19.4	.750 1.00 29.00 337 1.00 35.65 457 1.00 30.27 272 1.00 36.49
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3 ATOM 651 O PHE 239 48.151 29.521 19.4 ATOM 652 N SER 240 48.133 30.037 17.3	.750 1.00 29.00 337 1.00 35.65 457 1.00 30.27 272 1.00 36.49 3.359 1.00 36.37
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3 ATOM 651 O PHE 239 48.151 29.521 19.4 ATOM 652 N SER 240 48.133 30.037 17.3 45 ATOM 653 CA SER 240 46.936 30.866 17 ATOM 654 CB SER 240 46.622 31.466 15 ATOM 655 C SER 240 45.707 30.145 17.5	.750 1.00 29.00 337 1.00 35.65 457 1.00 30.27 272 1.00 36.49 359 1.00 36.37 .994 1.00 35.87
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3 ATOM 651 O PHE 239 48.151 29.521 19.4 ATOM 652 N SER 240 48.133 30.037 17.3 45 ATOM 653 CA SER 240 46.936 30.866 17 ATOM 654 CB SER 240 46.622 31.466 15 ATOM 655 C SER 240 45.707 30.145 17.9 ATOM 656 O SER 240 44.784 30.789 18.4	.750 1.00 29.00 337 1.00 35.65 457 1.00 30.27 272 1.00 36.49 359 1.00 36.37 .994 1.00 35.87 936 1.00 40.37 438 1.00 37.47
ATOM 649 CZ PHE 239 51.813 32.576 20 ATOM 650 C PHE 239 48.647 29.434 18.3 ATOM 651 O PHE 239 48.151 29.521 19.4 ATOM 652 N SER 240 48.133 30.037 17.3 45 ATOM 653 CA SER 240 46.936 30.866 17 ATOM 654 CB SER 240 46.622 31.466 15 ATOM 655 C SER 240 45.707 30.145 17.9	.750 1.00 29.00 337 1.00 35.65 457 1.00 30.27 272 1.00 36.49 3.359 1.00 36.37 .994 1.00 35.87 936 1.00 40.37 438 1.00 37.47 .889 1.00 43.00

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            660 CG GLU 241
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            678 CG PRO 243
     ATOM 679 C PRO 243
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	ATOM	710 NE2 GLN 247	46.439 32.185 20.729 1.00 35.19
	ATOM	710 NE2 GLN 247	51.190 33.196 24.575 1.00 27.51
	ATOM	711 C GLN 247 712 O GLN 247	52.377 33.261 24.256 1.00 28.70
5	ATOM		50.661 33.921 25.552 1.00 27.81
ر	ATOM	714 CA ILE 248	51.431 34.908 26.295 1.00 29.41
	ATOM	714 CA ILE 248 715 CB ILE 248	50,525 35.662 27.303 1.00 28.96
	ATOM	716 CG2 ILE 248	51.356 36.476 28.279 1.00 28.67
	ATOM	717 CG1 ILE 248	49.555 36.571 26.543 1.00 28.83
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	ATOM		52.356 33.177 27.732 1.00 26.07
	ATOM		53.413 32.474 28.454 1.00 27.37
1.5	ATOM	722 CA ILE 249	52.839 31.294 29.281 1.00 30.32
15	ATOM	723 CB ILE 249 724 CG2 ILE 249	53.958 30.425 29.840 1.00 31.29
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	ATOM		51.295 30.753 31.230 1.00 31.30
	ATOM	726 CD1 ILE 249	54.510 31.974 27.509 1.00 31.30
20	ATOM		55.701 32.100 27.808 1.00 29.59
20	ATOM	728 O ILE 249 729 N LEU 250	54.110 31.442 26.357 1.00 29.03
	ATOM	730 CA LEU 250	55.068 30.934 25.380 1.00 22.44
	ATOM	730 CA LEU 250 731 CB LEU 250	54.351 30.166 24.266 1.00 24.30
	ATOM	731 CB LEU 250 732 CG LEU 250	53.665 28.866 24.687 1.00 23.20
25	ATOM ATOM	732 CG LEU 250 733 CD1 LEU 250	52.951 28.273 23.502 1.00 20.36
23	ATOM	733 CD1 LEU 250 734 CD2 LEU 250	54.685 27.880 25.238 1.00 19.45
	ATOM	734 CD2 LEU 250	55.919 32.055 24.794 1.00 18.97
	ATOM	736 O LEU 250	57.133 31.903 24.648 1.00 18.37
	ATOM	737 N LEU 251	55.291 33.180 24.468 1.00 20.63
30	ATOM	737 N LEO 231 738 CA LEU 251	56.026 34.318 23.915 1.00 27.43
30	ATOM	739 CB LEU 251	55.065 35.412 23.449 1.00 22.92
	ATOM	740 CG LEU 251	54.364 35.093 22.128 1.00 24.72
	ATOM	740 CG LEO 251 741 CD1 LEU 251	53.342 36.167 21.821 1.00 32.13
	ATOM	741 CD1 LEU 251	55.389 34.981 21.009 1.00 22.46
35	ATOM	743 C LEU 251	57.026 34.875 24.930 1.00 27.23
23	ATOM	744 O LEU 251	58.202 35.078 24.614 1.00 26.48
	ATOM	745 N LYS 252	56.561 35.094 26.156 1.00 27.34
	ATOM	746 CA LYS 252	57.425 35.598 27.215 1.00 28.95
	ATOM	740 CA L13 252 747 CB LYS 252	56.649 35.715 28.527 1.00 32.89
40	ATOM	747 CB LTS 232 748 CG LYS 252	55.570 36.783 28.530 1.00 35.06
70	ATOM	748 CG L13 252 749 CD LYS 252	55.084 37.028 29.943 1.00 42.82
	ATOM	750 CE LYS 252	54.124 38.191 30.003 1.00 53.05
	ATOM	750 CE LTS 252 751 NZ LYS 252	53.677 38.451 31.398 1.00 64.03
	ATOM	751 NZ E13 232 752 C LYS 252	58.605 34.647 27.405 1.00 27.66
45	ATOM	752 C L1S 252 753 O LYS 252	59.734 35.076 27.646 1.00 33.16
73	ATOM	754 N GLY 253	58.344 33.357 27.243 1.00 24.50
	ATOM	755 CA GLY 253	59.386 32.364 27.402 1.00 22.33
	ATOM	756 C GLY 253	60.423 32.273 26.297 1.00 23.99
	ATOM	757 O GLY 253	61.589 32.016 26.581 1.00 30.77
50	ATOM	757 O GL 1 253 758 N CYS 254	60.041 32.526 25.049 1.00 22.66
50	ATOM	736 IV C13 234	00.041 32.320 23.047 1.00 22.00

PCT/US98/25296

# WO 99/26966

	ATOM	759 CA CYS 254	60.986 32.405 23.934 1.00 20.75
	ATOM	760 CB CYS 254	60.386 31.494 22.868 1.00 24.86
	ATOM	761 SG CYS 254	58.996 32.276 22.014 1.00 25.55
	ATOM	762 C CYS 254	61.399 33.702 23.242 1.00 23.79
5	<b>ATOM</b>	763 O CYS 254	62.262 33.685 22.357 1.00 22.18
	<b>ATOM</b>	764 N CYS 255	60.788 34.814 23.625 1.00 19.49
	<b>ATOM</b>	765 CA CYS 255	61.084 36.085 22.981 1.00 21.08
	<b>ATOM</b>	766 CB CYS 255	60.336 37.220 23.669 1.00 18.21
	<b>ATOM</b>	767 SG CYS 255	60.264 38.713 22.677 1.00 22.96
10	<b>ATOM</b>	768 C CYS 255	62.570 36.413 22.842 1.00 21.87
	ATOM	769 O CYS 255	63.050 36.641 21.729 1.00 22.23
	ATOM	770 N MET 256	63.310 36.397 23.947 1.00 20.82
	ATOM	771 CA MET 256	64.741 36.706 23.895 1.00 20.50
	<b>ATOM</b>	772 CB MET 256	65.322 36.801 25.312 1.00 22.50
15	ATOM	773 CG MET 256	66.808 37.139 25.354 1.00 16.67
	ATOM	774 SD MET 256	67.205 38.732 24.605 1.00 24.46
	ATOM	775 CE MET 256	69.027 38.764 24.791 1.00 19.21
	ATOM	776 C MET 256	65.510 35.667 23.072 1.00 18.38
	ATOM	777 O MET 256	66.401 36.005 22.293 1.00 17.68
20	ATOM	778 N GLU 257	65.149 34.404 23.248 1.00 20.33
	ATOM	779 CA GLU 257	65.779 33.308 22.526 1.00 21.08
	ATOM	780 CB GLU 257	65.148 31.982 22.943 1.00 22.28
	ATOM	781 CG GLU 257	65.374 31.640 24.411 1.00 34.68
	ATOM	782 CD GLU 257	64.515 30.486 24.907 1.00 43.20
25	ATOM	783 OE1 GLU 257	
	ATOM	784 OE2 GLU 257	
	ATOM	785 C GLU 257	65.650 33.503 21.018 1.00 19.26
	ATOM	786 O GLU 257	66.632 33.360 20.276 1.00 18.09
	ATOM	787 N ILE 258	64.446 33.850 20.566 1.00 16.30
30	ATOM	788 CA ILE 258	64.199 34.065 19.141 1.00 18.09
	ATOM	789 CB ILE 258	62.677 34.150 18.825 1.00 18.61
	ATOM	790 CG2 ILE 258	62.441 34.653 17.395 1.00 16.23
	ATOM	791 CG1 ILE 258	62.032 32.771 19.021 1.00 13.80 60.544 32.714 18.695 1.00 13.21
	ATOM	792 CD1 ILE 258	
35	ATOM	793 C ILE 258	64.948 35.297 18.638 1.00 20.12 65.605 35.242 17.593 1.00 19.17
	ATOM	794 O ILE 258 795 N MET 259	
	ATOM	795 N MET 259 796 CA MET 259	
	ATOM	790 CA MET 239 797 CB MET 259	65.249 38.772 19.941 1.00 18.80
40	ATOM	797 CB MET 239 798 CG MET 259	63.782 39.159 19.894 1.00 17.66
40	ATOM ATOM	798 CG MET 239 799 SD MET 259	63.457 40.748 20.678 1.00 25.77
	ATOM	800 CE MET 259	63.774 40.377 22.374 1.00 16.65
	ATOM	801 C MET 259	67.111 37.397 18.973 1.00 19.51
	ATOM	801 C MET 259 802 O MET 259	67.797 37.913 18.080 1.00 25.53
45	ATOM	802 O MET 239 803 N SER 260	67.797 37.913 18.080 1.00 25.55
43	ATOM	804 CA SER 260	69.056 36.324 19.947 1.00 16.90
	ATOM	804 CA SER 200 805 CB SER 260	69.434 35.631 21.251 1.00 15.56
	ATOM	806 OG SER 260	69.093 36.455 22.352 1.00 22.98
	ATOM	807 C SER 260	69.471 35.487 18.746 1.00 14.52
50	ATOM	808 O SER 260	70.496 35.761 18.129 1.00 22.82
-		545 C DEAC 200	

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            810 CA LEU 261
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            811 CB LEU 261
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            812 CG LEU 261
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            813 CD1 LEU 261
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            816 O LEU 261
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            823 CZ ARG 262
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            825 NH2 ARG 262
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            828 N ALA 263
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     ATOM
            838 N VAL 265
     ATOM
            839 CA VAL 265
                                72.380 34.226 12.306 1.00 28.98
            840 CB VAL 265
                                71.072 33.547 11.797 1.00 25.97
     ATOM
                                70.751 32.330 12.638 1.00 26.27
     ATOM
            841 CG1 VAL 265
                                69.907 34.527 11.797 1.00 26.64
     ATOM
            842 CG2 VAL 265
            843 C VAL 265
                               72.761 35.373 11.369 1.00 28.81
35
    ATOM
            844 O VAL 265
                               72.966 35.160 10.176 1.00 31.92
     ATOM
                               72.830 36.587 11.915 1.00 31.83
     ATOM
            845 N ARG 266
                                73.210 37.774 11.150 1.00 33.19
     ATOM
            846 CA ARG 266
                                72.141 38.861 11.258 1.00 31.67
     ATOM
            847 CB ARG 266
                                70.986 38.623 10.320 1.00 26.82
40
     ATOM
            848 CG ARG 266
                                69.913 39.668 10.454 1.00 33.95
     ATOM
            849 CD ARG 266
                                68.955 39.532 9.361 1.00 38.15
            850 NE ARG 266
     ATOM
                                67.688 39.927 9.410 1.00 37.39
     ATOM
            851 CZ ARG 266
                                 67.198 40.491 10.509 1.00 29.92
     ATOM
            852 NH1 ARG 266
                                 66.918 39.770 8.340 1.00 31.24
45
     ATOM
            853 NH2 ARG 266
            854 C ARG 266
                               74.565 38.307 11.604 1.00 36.31
     ATOM
                               74.821 39.516 11.575 1.00 38.56
     ATOM
            855 O ARG 266
            856 N TYR 267
                               75.416 37.393 12.056 1.00 34.21
     ATOM
                                76.755 37.733 12.502 1.00 35.24
            857 CA TYR 267
     ATOM
                                77.283 36.640 13.440 1.00 32.37
50
            858 CB TYR 267
     ATOM
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78.774 36.699 13.703 1.00 35.07
    ATOM
            859 CG TYR 267
            860 CD1 TYR 267
                                79.303 37.555 14.669 1.00 33.94
    ATOM
            861 CE1 TYR 267
                                80.677 37.609 14.905 1.00 36.60
    ATOM
                                79.658 35.894 12.979 1.00 34.68
            862 CD2 TYR 267
    ATOM
                                81.029 35.940 13.208 1.00 36.07
            863 CE2 TYR 267
    ATOM
                                81.533 36.797 14.170 1.00 37.14
            864 CZ TYR 267
    ATOM
                                82.889 36.835 14.396 1.00 41.52
            865 OH TYR 267
    ATOM
                               77.639 37.831 11.263 1.00 37.68
            866 C TYR 267
    ATOM
                               77.609 36.943 10.410 1.00 36.48
            867 O TYR 267
    ATOM
                               78,400 38,915 11.150 1.00 39.58
            868 N ASP 268
10
    ATOM
                               79.301 39.096 10.016 1.00 42.77
            869 CA ASP 268
    ATOM
                               79.170 40.511 9.434 1.00 44.38
            870 CB ASP 268
    ATOM
                                80.145 40.770 8.290 1.00 50.31
            871 CG ASP 268
    ATOM
            872 OD1 ASP 268
                                80.290 39.901 7.400 1.00 55.79
     ATOM
            873 OD2 ASP 268
                                80.773 41.847 8.280 1.00 50.24
     ATOM
15
                               80,737 38.846 10.466 1.00 42.51
     ATOM
            874 C ASP 268
                               81.305 39.645 11.208 1.00 42.75
    ATOM
            875 O ASP 268
                               81.346 37.733 10.020 1.00 44.56
            876 N PRO 269
     ATOM
                                80.770 36.697 9.146 1.00 42.66
            877 CD PRO 269
     ATOM
            878 CA PRO 269
                                82.725 37.395 10.393 1.00 45.98
20
     ATOM
                                82.991 36.111 9.607 1.00 44.04
            879 CB PRO 269
     ATOM
                                81.631 35.506 9.458 1.00 43.33
            880 CG PRO 269
     ATOM
                               83.710 38.492 10.004 1.00 50.31
     ATOM
            881 C PRO 269
                               84.630 38.800 10.761 1.00 49.83
            882 O PRO 269
     ATOM
                               83.486 39.100 8.840 1.00 53.62
            883 N ALA 270
25
     ATOM
                                84.348 40.165 8.329 1.00 54.54
            884 CA ALA 270
     ATOM
                                83.892 40.585 6.929 1.00 51.24
     ATOM
            885 CB ALA 270
                               84.449 41.389 9.248 1.00 55.69
            886 C ALA 270
     ATOM
            887. O ALA 270
                               85.488 42.045 9.294 1.00 57.92
     ATOM
                               83.384 41.685 9.989 1.00 54.71
            888 N SER 271
     ATOM
30
            889 CA SER 271
                                83.378 42.838 10.889 1.00 51.26
     ATOM
                                82.182 43.740 10.575 1.00 49.92
             890 CB SER 271
     ATOM
                                82.065 43.976 9.183 1.00 60.09
            891 OG SER 271
     ATOM
                               83.305 42.443 12.360 1.00 50.78
            892 C SER 271
     ATOM
                               83.482 43.288 13.240 1.00 52.11
            893 O SER 271
35
     ATOM
                               83.051 41.162 12.619 1.00 48.96
            894 N ASP 272
     ATOM
                                82.898 40.643 13.978 1.00 45.53
            895 CA ASP 272
     ATOM
             896 CB ASP 272
                                84.206 40.765 14.776 1.00 44.82
     ATOM
                                84.142 40.064 16.131 1.00 47.66
             897 CG ASP 272
     ATOM
             898 OD1 ASP 272
                                84.750 40.581 17.091 1.00 48.64
40
     ATOM
             899 OD2 ASP 272
                                83.495 38.999 16.238 1.00 43.85
     ATOM
                               81.765 41.437 14.636 1.00 44.46
             900 C ASP 272
     ATOM
             901 O ASP 272
                               81.904 41.958 15.747 1.00 42.41
     ATOM
                               80.652 41.551 13.915 1.00 39.79
             902 N THR 273
     ATOM
                                79.492 42.282 14.401 1.00 38.82
             903 CA THR 273
45
     ATOM
                                79.334 43.648 13.670 1.00 39.73
     ATOM
             904 CB THR 273
                                 79.288 43.439 12.254 1.00 39.36
             905 OG1 THR 273
     ATOM
                                 80.496 44.578 13.991 1.00 41.31
             906 CG2 THR 273
     ATOM
                               78.203 41.485 14.211 1.00 38.36
     ATOM
            907 C THR 273
            908 O THR 273
                               78.151 40.546 13.408 1.00 33.79
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     ATOM
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ATOM 911 CB LEU 274 75.369 41.212 14.912 1.00 34.49 ATOM 912 CG LEU 274 75.948 39.651 17.069 1.00 30.37 ATOM 913 CD1 LEU 274 75.948 39.651 17.069 1.00 32.97 75.948 39.651 17.069 1.00 32.97 75.948 39.651 17.069 1.00 32.97 75.949 39.593 18.440 1.00 28.23 ATOM 914 CD2 LEU 274 75.546 42.289 14.352 1.00 35.35 ATOM 916 O LEU 274 75.956 42.289 14.352 1.00 35.35 ATOM 917 N THR 275 73.942 41.890 13.599 1.00 34.05 ATOM 919 CB THR 275 73.942 41.890 13.599 1.00 34.05 ATOM 919 CB THR 275 73.942 41.890 13.599 1.00 34.05 ATOM 920 OG1 THR 275 71.699 42.746 13.793 1.00 30.94 ATOM 921 CG2 THR 275 71.699 42.746 13.793 1.00 30.94 ATOM 922 C THR 275 71.699 42.746 13.793 1.00 30.92 ATOM 924 N LEU 276 70.051 43.868 15.192 1.00 27.78 ATOM 925 CA LEU 276 70.051 43.868 15.192 1.00 27.78 ATOM 927 CG LEU 276 70.051 43.868 15.192 1.00 27.78 ATOM 929 CD2 LEU 276 71.383 44.532 17.373 1.00 25.89 ATOM 931 O LEU 276 68.930 44.376 14.296 1.00 27.27 ATOM 932 N SER 277 66.697 43.957 13.366 1.00 20.79 ATOM 932 N SER 277 67.854 43.598 14.187 1.00 25.97 ATOM 933 CA SER 277 66.697 43.957 13.366 1.00 20.79 ATOM 938 N GLY 278 ATOM 936 C SER 277 67.051 43.863 13.672 1.00 20.079 ATOM 937 O SER 277 66.697 43.957 13.366 1.00 20.79 ATOM 938 N GLY 278 ATOM 936 C SER 277 67.054 44.399 11.181 1.00 25.97 ATOM 937 O SER 277 66.697 43.957 13.366 1.00 28.63 ATOM 940 C GLY 278 ATOM 940 C GLU 279 ATOM 950 C BLU 279 ATOM 951 N MET 280 ATOM 952 CA MET 280 ATOM 955 CB MET 280 ATOM 957 C MET 280 ATOM 958 O M		ATOM	909 N LEU 274	77.187 41.835 14.995 1.00 36.91
ATOM 911 CB LEU 274 75.342 40.822 16.297 1.00 30.37 ATOM 912 CG LEU 274 75.297 39.593 18.440 1.00 28.23 75.749 38.341 16.318 1.00 26.86 ATOM 915 C LEU 274 74.75.94 38.341 16.318 1.00 26.86 ATOM 916 O LEU 274 75.749 38.341 16.318 1.00 26.86 ATOM 917 N THR 275 74.956 42.289 14.352 1.00 35.35 74.70M 917 N THR 275 73.942 41.890 13.599 1.00 34.05 ATOM 919 CB THR 275 73.942 41.890 13.599 1.00 34.05 ATOM 920 OG1 THR 275 74.108 42.590 10.909 1.00 39.50 ATOM 921 CG2 THR 275 74.108 42.590 10.909 1.00 39.50 72.064 43.851 10.952 1.00 30.94 ATOM 922 C THR 275 71.100 41.670 13.845 1.00 36.53 ATOM 924 N LEU 276 70.051 43.868 15.192 1.00 28.10 ATOM 925 CA LEU 276 70.051 43.868 15.192 1.00 22.78 ATOM 926 CB LEU 276 70.051 43.868 15.192 1.00 22.78 ATOM 927 CG LEU 276 70.051 43.868 15.192 1.00 22.78 ATOM 928 CD1 LEU 276 70.051 43.868 15.192 1.00 20.70 ATOM 929 CD2 LEU 276 70.051 43.868 15.192 1.00 20.70 ATOM 932 N SER 277 70.205 44.780 16.420 1.00 22.51 ATOM 931 O LEU 276 69.068 45.430 13.672 1.00 20.79 ATOM 932 N SER 277 66.597 43.957 13.366 1.00 22.07 71.456 43.099 17.782 1.00 20.79 ATOM 936 C SER 277 66.597 43.957 13.366 1.00 22.65 ATOM 937 O SER 277 65.561 44.905 15.290 1.00 22.65 ATOM 940 C GLY 278 ATOM 941 O GLY 278 ATOM 942 N GLU 279 ATOM 942 N GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 947 OE1 GLU 279 ATOM 949 C GLU 279 ATOM 949 C GLU 279 ATOM 940 C GLU 279 ATOM 950 O GLU 279 ATOM 951 N MET 280 ATOM 952 CA MET 280 ATOM 955 CB MET 280 ATOM 955 CB MET 280 ATOM 955 CB MET 280 ATOM 957 C MET 280 ATOM 955 C MET 280 ATOM 957 C MET 280 ATOM 955 C MET 28			910 CA LEU 274	75.869 41.212 14.912 1.00 34.49
5 ATOM 913 CDI LEU 274 75.948 39.651 17.069 1.00 32.97 ATOM 914 CD2 LEU 274 75.297 39.593 18.440 1.00 28.23 75.749 38.341 16.318 1.00 26.86 1.00 26.251 75.749 38.341 16.318 1.00 26.86 1.00 37.47 75.97 39.593 18.440 1.00 28.23 75.749 38.341 16.318 1.00 26.86 1.00 37.47 75.97 39.593 18.440 1.00 28.23 75.749 38.341 16.318 1.00 26.86 1.00 37.47 75.97 39.593 18.440 1.00 28.23 75.74 30.00 37.47 75.75 30.00 37.47 75.75 30.00 37.47 75.75 30.00 37.47 75.75 30.00 42.868 13.052 1.00 37.47 75.75 30.00 42.868 13.052 1.00 34.05 72.00 42.868 13.052 1.00 34.05 72.00 42.868 13.052 1.00 34.05 72.00 42.868 13.052 1.00 32.62 72.00 42.868 13.052 1.00 32.62 72.00 42.868 13.052 1.00 32.62 72.00 42.868 13.052 1.00 32.62 72.00 42.868 13.052 1.00 30.94 75.00 92.00 10.00 92.00 32.62 72.00 42.868 13.052 1.00 30.94 75.00 92.00				75.342 40.822 16.297 1.00 30.37
5         ATOM         913         CD1 LEU         274         75.297         39.593         18.440         1.00 28.23           ATOM         914         CD2 LEU         274         75.749         38.341         16.318         1.00 26.86           ATOM         916         O LEU         274         74.956         42.289         14.352         1.00 35.37           ATOM         917         N THR         275         73.942         41.890         13.599         1.00 37.47           ATOM         919         CB THR         275         73.020         42.868         13.052         1.00 32.62           ATOM         910         CB THR         275         73.020         42.868         13.052         1.00 30.94           ATOM         920         OG1 THR         275         73.020         42.868         13.052         1.00 30.94           ATOM         920         OG1 THR         275         71.069         42.746         13.793         1.00 30.94           15         ATOM         921         CG LEU         276         70.025         44.780         13.943         1.00 30.94           15         ATOM         925         CA LEU         276         70.051		ATOM	912 CG LEU 274	75.948 39.651 17.069 1.00 32.97
ATOM 914 CD2 LEU 274 ATOM 915 C LEU 274 ATOM 916 O LEU 274 ATOM 917 N THR 275 ATOM 918 CA THR 275 ATOM 919 CB THR 275 ATOM 919 CB THR 275 ATOM 919 CG THR 275 ATOM 921 CG2 THR 275 ATOM 922 C THR 275 ATOM 923 O THR 275 ATOM 924 N LEU 276 ATOM 925 CA LEU 276 ATOM 927 CG LEU 276 ATOM 930 C LEU 276 ATOM 931 O LEU 276 ATOM 931 C LEU 276 ATOM 932 C THR 275 ATOM 926 CB LEU 276 ATOM 927 CG LEU 276 ATOM 930 C LEU 276 ATOM 931 C LEU 276 ATOM 931 C LEU 276 ATOM 932 N SER 277 ATOM 934 CB SER 277 ATOM 935 CG SER 277 ATOM 936 C SER 277 ATOM 937 O SER 277 ATOM 938 N GLY 278 ATOM 939 CA GLY 278 ATOM 940 C GLY 278 ATOM 940 C GLY 278 ATOM 941 C GLY 276 ATOM 942 N GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLU 279 ATOM 947 OEI GLU 279 ATOM 948 CEZ GLU 279 ATOM 949 C GLU 279 ATOM 940 C GLU 279 ATOM 940 C GLU 279 ATOM 941 O GLY 278 ATOM 942 N GLU 279 ATOM 943 CA GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLU 279 ATOM 947 OEI GLU 279 ATOM 948 OEZ GLU 279 ATOM 949 C GLU 279 ATOM 940 C GLU 279 ATOM 941 O GLY 278 ATOM 942 N GLU 279 ATOM 943 CA GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLU 279 ATOM 947 OEI GLU 279 ATOM 948 OEZ GLU 279 ATOM 949 C GLU 279 ATOM 940 C GLU 279 ATOM 941 O GLU 279 ATOM 942 N GLU 279 ATOM 943 CA GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLU 279 ATOM 947 OEI GLU 279 ATOM 948 OEZ GLU 279 ATOM 949 C GLU 279 ATOM 940 CG GLU 279 ATOM 940 CG GLU 279 ATOM 941 OEI GLU 279 ATOM 942 CA GLU 279 ATOM 943 CA GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLU 279 ATOM 947 OEI GLU 279 ATOM 948 OEZ GLU 279 ATOM 949 C GLU 279 ATOM 950 O GLU 279 ATOM 950 O GLU 279 ATOM 950 CA MET 280 ATOM 951 N MET 280 ATOM 952 CA MET 280 ATOM 953 CB MET 280 ATOM 955 CM MET 280 ATOM 956 C MET 280 ATOM 957 C MET 280 ATOM 95	5			
ATOM 916 C LEU 274 74,956 42.289 14.352 1.00 35.35 ATOM 916 O LEU 274 73,942 41.890 13.599 1.00 34.05 73.020 42.868 13.052 1.00 32.62 ATOM 919 CB THR 275 72.824 42.674 11.542 1.00 35.14 ATOM 920 OG1 THR 275 72.824 42.674 11.542 1.00 35.14 ATOM 921 CG2 THR 275 72.064 43.851 10.952 1.00 30.94 ATOM 922 C THR 275 71.699 42.746 13.793 1.00 30.92 71.00 41.670 13.845 1.00 36.53 ATOM 925 CA LEU 276 ATOM 925 CA LEU 276 ATOM 926 CB LEU 276 ATOM 927 CG LEU 276 ATOM 929 CD2 LEU 276 ATOM 930 C LEU 276 ATOM 931 O LEU 276 ATOM 931 C LEU 276 ATOM 932 N SER 277 ATOM 931 C LEU 276 ATOM 932 N SER 277 ATOM 934 CB SER 277 ATOM 935 OG SER 277 ATOM 936 C SER 277 ATOM 937 O SER 277 ATOM 938 N GLY 278 ATOM 939 CA GLY 278 ATOM 939 CA GLY 278 ATOM 939 CA GLY 278 ATOM 930 C LEU 276 ATOM 931 O LEU 276 ATOM 934 CB SER 277 ATOM 935 OG SER 277 ATOM 936 C SER 277 ATOM 937 O SER 277 ATOM 937 O SER 277 ATOM 938 N GLY 278 ATOM 940 C GLY 278 ATOM 940 C GLY 278 ATOM 941 O GLY 278 ATOM 942 N GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLU 279 ATOM 947 OE1 GLU 279 ATOM 948 OE2 GLU 279 ATOM 949 C GLU 279 ATOM 940 C GLU 279 ATOM 950 O GLU 279 ATOM 951 N MET 280 ATOM 955 CM MET 280 ATOM 955 CM MET 280 ATOM 955 C MET 280 ATOM 957 C MET 2	•			
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ATOM 917 N THR 275 73.942 41.890 13.599 1.00 34.05 ATOM 918 CA THR 275 73.020 42.868 13.052 1.00 32.62 ATOM 920 OG1 THR 275 74.108 42.590 10.909 1.00 39.50 ATOM 921 CG2 THR 275 74.108 42.590 10.909 1.00 39.50 ATOM 922 C THR 275 71.699 42.746 13.793 1.00 30.94 ATOM 924 N LEU 276 71.291 43.835 14.434 1.00 28.10 ATOM 925 CA LEU 276 70.205 44.780 16.420 1.00 22.51 ATOM 926 CB LEU 276 70.205 44.780 16.420 1.00 22.51 ATOM 928 CD1 LEU 276 71.295 43.608 1.00 20.70 ATOM 929 CD2 LEU 276 71.295 43.408 18.608 1.00 20.70 ATOM 920 CD2 LEU 276 71.295 43.408 18.608 1.00 20.70 ATOM 920 CD2 LEU 276 71.295 43.409 13.672 1.00 20.79 ATOM 930 C LEU 276 68.930 44.376 14.296 1.00 27.27 ATOM 931 O LEU 276 67.854 43.598 14.187 1.00 25.97 ATOM 932 N SER 277 66.697 43.957 13.366 1.00 22.65 ATOM 933 CA SER 277 65.561 44.905 15.290 1.00 22.65 67.864 44.995 15.290 1.00 22.65 ATOM 938 N GLY 278 68.688 43.597 11.181 1.00 28.52 ATOM 940 C GLY 278 ATOM 940 C GLY 278 ATOM 940 C GLY 278 ATOM 944 CB GLU 279 ATOM 944 CB GLU 279 ATOM 945 CG GLU 279 ATOM 946 CD GLY 278 ATOM 947 OE1 GLU 279 ATOM 948 OE2 GLU 279 ATOM 948 OE2 GLU 279 ATOM 949 C GLU 279 ATOM 940 C GLU 279 ATOM 94				
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ATOM 944 CB GLU 279 68.314 48.312 10.515 1.00 44.28 ATOM 945 CG GLU 279 67.703 48.322 11.908 1.00 52.30 ATOM 946 CD GLU 279 66.440 49.159 12.001 1.00 60.23 ATOM 947 OE1 GLU 279 66.398 50.074 12.853 1.00 63.06 40 ATOM 948 OE2 GLU 279 65.485 48.894 11.238 1.00 65.67 ATOM 949 C GLU 279 70.700 48.038 11.216 1.00 42.40 ATOM 950 O GLU 279 71.330 49.001 10.787 1.00 43.89 ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86 ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80 45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77		ATOM		
ATOM 945 CG GLU 279 67.703 48.322 11.908 1.00 52.30 ATOM 946 CD GLU 279 66.440 49.159 12.001 1.00 60.23 ATOM 947 OE1 GLU 279 66.398 50.074 12.853 1.00 63.06  40 ATOM 948 OE2 GLU 279 65.485 48.894 11.238 1.00 65.67 ATOM 949 C GLU 279 70.700 48.038 11.216 1.00 42.40 ATOM 950 O GLU 279 71.330 49.001 10.787 1.00 43.89 ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86 ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80  45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77	35	ATOM		
ATOM 946 CD GLU 279 66.440 49.159 12.001 1.00 60.23 ATOM 947 OE1 GLU 279 66.398 50.074 12.853 1.00 63.06  40 ATOM 948 OE2 GLU 279 65.485 48.894 11.238 1.00 65.67 ATOM 949 C GLU 279 70.700 48.038 11.216 1.00 42.40 ATOM 950 O GLU 279 71.330 49.001 10.787 1.00 43.89 ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86 ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80  45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
ATOM 947 OE1 GLU 279 66.398 50.074 12.853 1.00 63.06  40 ATOM 948 OE2 GLU 279 65.485 48.894 11.238 1.00 65.67  ATOM 949 C GLU 279 70.700 48.038 11.216 1.00 42.40  ATOM 950 O GLU 279 71.330 49.001 10.787 1.00 43.89  ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86  ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80  45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25  ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64  ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02  ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56  ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				0,
40       ATOM       948       OE2 GLU       279       65.485       48.894       11.238       1.00       65.67         ATOM       949       C       GLU       279       70.700       48.038       11.216       1.00       42.40         ATOM       950       O       GLU       279       71.330       49.001       10.787       1.00       43.89         ATOM       951       N       MET       280       70.977       47.472       12.388       1.00       40.86         ATOM       952       CA       MET       280       72.027       48.009       13.248       1.00       32.80         45       ATOM       953       CB       MET       280       71.435       48.415       14.603       1.00       32.80         45       ATOM       954       CG       MET       280       72.384       49.193       15.506       1.00       31.64         ATOM       955       SD       MET       280       71.830       49.235       17.232       1.00       34.02         ATOM       956       CE       MET       280       70.566       50.495       17.197       1.00       26.56				
ATOM 949 C GLU 279 70.700 48.038 11.216 1.00 42.40 ATOM 950 O GLU 279 71.330 49.001 10.787 1.00 43.89 ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86 ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80  45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
ATOM 950 O GLU 279 71.330 49.001 10.787 1.00 43.89 ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86 ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80 45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77	40			
ATOM 951 N MET 280 70.977 47.472 12.388 1.00 40.86 ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80 45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
ATOM 952 CA MET 280 72.027 48.009 13.248 1.00 32.80 45 ATOM 953 CB MET 280 71.435 48.415 14.603 1.00 29.25 ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
45       ATOM       953       CB       MET       280       71.435       48.415       14.603       1.00       29.25         ATOM       954       CG       MET       280       72.384       49.193       15.506       1.00       31.64         ATOM       955       SD       MET       280       71.830       49.235       17.232       1.00       34.02         ATOM       956       CE       MET       280       70.566       50.495       17.197       1.00       26.56         ATOM       957       C       MET       280       73.172       47.033       13.465       1.00       32.77				
ATOM 954 CG MET 280 72.384 49.193 15.506 1.00 31.64 ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
ATOM 955 SD MET 280 71.830 49.235 17.232 1.00 34.02 ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77	45			
ATOM 956 CE MET 280 70.566 50.495 17.197 1.00 26.56 ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
ATOM 957 C MET 280 73.172 47.033 13.465 1.00 32.77				
				/U.366 3U.493 17.197 1.00 26.36
30 AIUM 938 U MEI 280 /2.983 43.9/1 14.038 1.00 34.61	<b>5</b> 0			
	50	ATOM	958 O MEI 280	12.983 43.9/1 14.038 1.00 34.61

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74.351 47.375 12.959 1.00 31.87
            959 N ALA 281
     ATOM
                               75.523 46.526 13.147 1.00 34.71
            960 CA ALA 281
     ATOM
                               76.519 46.727 12.023 1.00 34.42
    ATOM
            961 CB ALA 281
                              76.125 46.950 14.482 1.00 36.76
            962 C ALA 281
    ATOM
                               76.416 48.129 14.693 1.00 34.59
            963 O ALA 281
    ATOM
                               76.275 45.993 15.390 1.00 37.16
           964 N VAL 282
    ATOM
                               76.798 46.263 16.721 1.00 37.83
           965 CA VAL 282
    ATOM
           966 CB VAL 282
                               75.692 46.023 17.780 1.00 37.58
    ATOM
           967 CG1 VAL 282
                                76.219 46.271 19.175 1.00 48.99
    ATOM
                                74.514 46.939 17.514 1.00 43.59
10
    ATOM
            968 CG2 VAL 282
                               78.017 45.400 17.046 1.00 39.04
     ATOM
            969 C VAL 282
                               78.081 44.230 16.660 1.00 39.16
     ATOM
            970 O VAL 282
     ATOM
            971 N LYS 283
                              78.989 45.993 17.735 1.00 38.75
            972 CA LYS 283
                               80.205 45.287 18.136 1.00 42.18
     ATOM
                               81.428 46.208 18.045 1.00 47.46
            973 CB LYS 283
15
     ATOM
            974 CG LYS 283
                               81.803 46.617 16.632 1.00 51.71
    ATOM
                               83.092 47.416 16.618 1.00 59.26
            975 CD LYS 283
     ATOM
                               83.481 47.813 15.202 1.00 62.52
     ATOM
            976 CE LYS 283
           977 NZ LYS 283
                               82,492 48,742 14,588 1,00 66,27
     ATOM
                              80.075 44.746 19.559 1.00 38.78
           978 C LYS 283
20
     ATOM
           979 O LYS 283
                              79.283 45.257 20.356 1.00 40.63
     ATOM
            980 N ARG 284
                               80.900 43.753 19.881 1.00 36.01
     ATOM
                                80.908 43.104 21.189 1.00 38.62
            981 CA ARG 284
     ATOM
                                82.150 42.224 21.327 1.00 38.83
            982 CB ARG 284
     ATOM
                                82.220 41.091 20.333 1.00 41.87
25
     ATOM
            983 CG ARG 284
                                83.521 40.335 20.451 1.00 39.60
     ATOM
            984 CD ARG 284
                                83.506 39.120 19.644 1.00 45.18
            985 NE ARG 284
     ATOM
            986 CZ ARG 284
                                83.259 37.905 20.128 1.00 44.79
     ATOM
                                83.005 37.739 21.421 1.00 41.84
            987 NH1 ARG 284
     ATOM
                                83,271 36,852 19.319 1.00 42.27
            988 NH2 ARG 284
     ATOM
30
                               80.829 44.051 22.385 1.00 41.18
            989 C ARG 284
     ATOM
                               79.995 43.867 23.274 1.00 44.38
            990 O ARG 284
     ATOM
                               81.703 45.052 22.416 1.00 38.71
     ATOM 991 N GLU 285
                               81.724 46.002 23.525 1.00 37.18
     ATOM 992 CA GLU 285
     ATOM 993 CB GLU 285
                                82.950 46.906 23.422 1.00 36.65
35
                               80.444 46.838 23.614 1.00 35.71
     ATOM 994 C GLU 285
                               79.921 47.074 24.704 1.00 33.00
            995 O GLU 285
     ATOM
                               79.920 47.245 22.463 1.00 32.01
            996 N GLN 286
     ATOM
                                78.714 48.061 22.425 1.00 32.31
            997 CA GLN 286
     ATOM
                                78.440 48.525 20.997 1.00 38.24
            998 CB GLN 286
40
     ATOM
                                79.565 49.352 20.392 1.00 42.42
     ATOM
            999 CG GLN 286
                                79.277 49.761 18.964 1.00 44.79
     ATOM 1000 CD GLN 286
     ATOM 1001 OE1 GLN 286
                                 79.103 48.910 18.089 1.00 42.21
                                79.215 51.063 18.719 1.00 47.53
     ATOM 1002 NE2 GLN 286
                               77.484 47.355 23.002 1.00 33.08
45
     ATOM 1003 C GLN 286
                                76.770 47.929 23.827 1.00 30.95
     ATOM 1004 O GLN 286
                               77.245 46.114 22.579 1.00 31.49
     ATOM 1005 N LEU 287
                                76.095 45.350 23.068 1.00 31.01
     ATOM 1006 CA LEU 287
     ATOM 1007 CB LEU 287
                                75.892 44.073 22.242 1.00 24.63
                                74.498 43.780 21.661 1.00 27.34
     ATOM 1008 CG LEU 287
50
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ATOM 1009 CD1 LEU 287
                                74.382 42.282 21.359 1.00 20.50
                                 73.393 44.205 22.616 1.00 14.41
    ATOM 1010 CD2 LEU 287
                               76.298 44.986 24.538 1.00 32.80
     ATOM 1011 C LEU 287
                               75.351 45.014 25.334 1.00 32.10
     ATOM 1012 O LEU 287
    ATOM 1013 N LYS 288
                               77.536 44.641 24.885 1.00 32.54
                                77.897 44.280 26.251 1.00 30.70
    ATOM 1014 CA LYS 288
                                79.376 43.893 26.315 1.00 31.24
    ATOM 1015 CB LYS 288
    ATOM 1016 CG LYS 288
                                79.834 43.382 27.662 1.00 34.69
    ATOM 1017 CD LYS 288
                                81.227 42.784 27.574 1.00 37.69
    ATOM 1018 CE LYS 288
                                81.638 42.177 28.904 1.00 42.86
10
    ATOM 1019 NZ LYS 288
                                82.883 41.369 28.786 1.00 49.63
                               77.611 45.448 27.189 1.00 28.74
     ATOM 1020 C LYS 288
    ATOM 1021 O LYS 288
                               76.827 45.319 28.129 1.00 34.45
    ATOM 1022 N ASN 289
                               78.190 46.602 26.882 1.00 26.57
    ATOM 1023 CA ASN 289
                                78.011 47.803 27.691 1.00 30.84
15
    ATOM 1024 CB ASN 289
                                79.012 48.879 27.274 1.00 26.04
    ATOM 1025 CG ASN 289
                                80.437 48.485 27.570 1.00 35.16
     ATOM 1026 OD1 ASN 289
                                 80.700 47.718 28.499 1.00 42.54
    ATOM 1027 ND2 ASN 289
                                 81.371 48.998 26.784 1.00 32.82
                               76.602 48.371 27.620 1.00 35.05
    ATOM 1028 C ASN 289
20
     ATOM 1029 O ASN 289
                                76.154 49.039 28.550 1.00 36.94
     ATOM 1030 N GLY 290
                                75.909 48.113 26.515 1.00 32.43
     ATOM 1031 CA GLY 290
                                74.556 48.614 26.345 1.00 28.66
     ATOM 1032 C GLY 290
                               73.525 48.024 27.289 1.00 28.48
     ATOM 1033 O GLY 290
                                72.377 48.467 27.308 1.00 28.17
25
     ATOM 1034 N GLY 291
                                73.908 47.002 28.047 1.00 28.66
                                72.969 46.408 28.980 1.00 29.19
     ATOM 1035 CA GLY 291
     ATOM 1036 C GLY 291
ATOM 1037 O GLY 291
                                72.976 44.894 29.075 1.00 29.76
                                72.595 44.340 30.105 1.00 34.44
     ATOM 1038 N LEU 292
                                73.399 44.213 28.017 1.00 29.69
30
                                73.410 42.755 28.036 1.00 30.64
     ATOM 1039 CA LEU 292
                                73.421 42.194 26.611 1.00 27.07
     ATOM 1040 CB LEU 292
                                72.113 42.348 25.833 1.00 23.27
     ATOM 1041 CG LEU 292
     ATOM 1042 CD1 LEU 292
                                 72.202 41.580 24.532 1.00 22.24
                                 70.950 41.827 26.661 1.00 23.80
     ATOM 1043 CD2 LEU 292
35
                                74.530 42.125 28.861 1.00 29.22
     ATOM 1044 C LEU 292
     ATOM 1045 O LEU 292
                                74.365 41.033 29.404 1.00 31.02
                                75.671 42.800 28.945 1.00 30.26
     ATOM 1046 N GLY 293
     ATOM 1047 CA GLY 293
                                76,788 42,259 29,700 1.00 28,37
                                77.307 40.995 29.040 1.00 29.85
     ATOM 1048 C GLY 293
40
     ATOM 1049 O GLY 293
ATOM 1050 N VAL 294
                                77.460 40.951 27.820 1.00 32.37
                                77.537 39.953 29.832 1.00 30.08
     ATOM 1051 CA VAL 294
                                 78.041 38.687 29.308 1.00 31.62
     ATOM 1052 CB VAL 294
                                 78.466 37.716 30.442 1.00 29.11
                                 79.649 38.292 31.191 1.00 31.37
     ATOM 1053 CG1 VAL 294
45
                                 77.304 37.443 31.396 1.00 26.69
     ATOM 1054 CG2 VAL 294
                                77.079 37.978 28.351 1.00 32.81
     ATOM 1055 C VAL 294
     ATOM 1056 O VAL 294
                                77.496 37.095 27.591 1.00 33.00
                                75.801 38.356 28.380 1.00 30.45
     ATOM 1057 N VAL 295
                                74.814 37.752 27.487 1.00 28.02
     ATOM 1058 CA VAL 295
50
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	ATOM	1059 CB VAL 295	73.378 38.232 27.793 1.00 29.96
	ATOM	1060 CG1 VAL 295	72.380 37.575 26.838 1.00 22.55
	ATOM	1061 CG2 VAL 295	73.016 37.903 29.232 1.00 20.10
	ATOM	1062 C VAL 295	75.203 38.115 26.057 1.00 29.90
5	ATOM	1063 O VAL 295	75.047 37.312 25.140 1.00 34.47
J	ATOM	1064 N SER 296	75.762 39.309 25.886 1.00 29.11
	ATOM	1065 CA SER 296	76.215 39.771 24.581 1.00 30.96
	ATOM	1066 CB SER 296	76.785 41.184 24.702 1.00 27.26
	ATOM	1067 OG SER 296	77.300 41.648 23.469 1.00 22.93
10	ATOM	1068 C SER 296	77.294 38.811 24.080 1.00 36.41
10	ATOM	1069 O SER 296	77.238 38.341 22.939 1.00 38.84
	ATOM	1070 N ASP 297	78.254 38.501 24.954 1.00 35.29
	ATOM	1070 N ASP 297	79.346 37.585 24.629 1.00 32.14
			80.245 37.356 25.851 1.00 36.57
1.5	ATOM		80.958 38.616 26.307 1.00 41.75
15	ATOM	=	81.492 39.352 25.447 1.00 45.45
	ATOM	1074 OD1 ASP 297	80.999 38.861 27.532 1.00 45.15
	ATOM	1075 OD2 ASP 297	78.768 36.249 24.191 1.00 29.61
	ATOM	1076 C ASP 297	79.242 35.644 23.231 1.00 32.90
20	ATOM	1077 O ASP 297	77.738 35.804 24.903 1.00 27.85
20	ATOM	1078 N ALA 298	77.071 34.544 24.608 1.00 27.89
	ATOM	1079 CA ALA 298	
	ATOM	1080 CB ALA 298	
	ATOM	1081 C ALA 298	76.462 34.539 23.202 1.00 28.26
0.5	ATOM	1082 O ALA 298	76.648 33.579 22.446 1.00 30.19
25	ATOM	1083 N ILE 299	75.744 35.606 22.853 1.00 25.20
	ATOM	1084 CA ILE 299	75.119 35.708 21.537 1.00 23.46
	ATOM	1085 CB ILE 299	74.200 36.944 21.427 1.00 21.63
	ATOM	1086 CG2 ILE 299	73.491 36.946 20.078 1.00 22.20
	ATOM	1087 CG1 ILE 299	73.145 36.914 22.536 1.00 19.79
30	ATOM	1088 CD1 ILE 299	72.245 38.139 22.578 1.00 18.33
	ATOM	1089 C ILE 299	76.181 35.752 20.444 1.00 26.28
	ATOM	1090 O ILE 299	76.043 35.095 19.414 1.00 31.72
	ATOM	1091 N PHE 300	77.247 36.512 20.675 1.00 29.35
	ATOM	1092 CA PHE 300	78.338 36.613 19.709 1.00 29.01
35	ATOM	1093 CB PHE 300	79.386 37.622 20.182 1.00 29.53
	ATOM	1094 CG PHE 300	79.239 38.978 19.562 1.00 27.60
	ATOM	1095 CD1 PHE 300	78.481 39.964 20.179 1.00 24.86
	ATOM	1096 CD2 PHE 300	79.853 39.266 18.350 1.00 27.39
	ATOM	1097 CE1 PHE 300	78.337 41.218 19.597 1.00 25.66
40	ATOM	1098 CE2 PHE 300	79.715 40.518 17.761 1.00 25.97
	ATOM	1099 CZ PHE 300	78.956 41.495 18.384 1.00 21.03
	ATOM	1100 C PHE 300	78.988 35.248 19.496 1.00 30.34
	. ATOM	1101 O PHE 300	79.309 34.873 18.367 1.00 29.35
	ATOM	1102 N GLU 301	79.181 34.507 20.582 1.00 31.04
45	ATOM	1103 CA GLU 301	79.775 33.178 20.499 1.00 33.60
	ATOM	1104 CB GLU 301	80.012 32.607 21.898 1.00 31.64
	ATOM	1105 C GLU 301	78.851 32.265 19.696 1.00 33.90
	<b>ATOM</b>	1106 O GLU 301	79.315 31.473 18.872 1.00 33.33
	ATOM	1107 N LEU 302	77.546 32.386 19.935 1.00 31.13
50	ATOM	1108 CA LEU 302	76.556 31.581 19.227 1.00 27.57

	ATOM	1109 CB LEU 302	75.150 31.842 19.776 1.00 25.24
	ATOM	1110 CG LEU 302	73.994 31.131 19.059 1.00 28.59
	ATOM	1111 CD1 LEU 302	74.066 29.634 19.299 1.00 25.52
	ATOM	1112 CD2 LEU 302	72.660 31.682 19.532 1.00 19.30
5	ATOM	1113 C LEU 302	76.601 31.904 17.739 1.00 26.80
,	ATOM	1114 O LEU 302	76.682 31.003 16.904 1.00 27.81
	ATOM	1115 N GLY 303	76.576 33.195 17.416 1.00 26.47
	ATOM	1116 CA GLY 303	76.611 33.624 16.030 1.00 26.99
	ATOM	1117 C GLY 303	77.845 33.133 15.295 1.00 33.46
10	ATOM	1118 O GLY 303	77.757 32.646 14.164 1.00 32.33
10	ATOM	1119 N LYS 304	78.994 33.232 15.956 1.00 34.63
	ATOM	1120 CA LYS 304	80.269 32.813 15.383 1.00 36.20
	ATOM	1121 CB LYS 304	81.399 33.115 16.372 1.00 41.96
		1122 CG LYS 304	82.779 33.179 15.757 1.00 47.05
1.5	ATOM	1122 CG L13 304 1123 CD LYS 304	83.800 33.610 16.796 1.00 59.47
15	ATOM	1124 CE LYS 304	85.179 33.791 16.181 1.00 65.89
	ATOM		85.182 34.863 15.144 1.00 71.01
	ATOM	1125 NZ LYS 304 1126 C LYS 304	80.276 31.332 14.992 1.00 33.17
	ATOM		80.752 30.974 13.913 1.00 34.44
20	ATOM		79.739 30.482 15.861 1.00 31.40
20	ATOM	1128 N SER 305 1129 CA SER 305	79.687 29.048 15.594 1.00 33.10
	ATOM	1129 CA SER 305 1130 CB SER 305	79.513 28.266 16.900 1.00 34.10
	ATOM	1130 CB SER 303	78.391 28.727 17.633 1.00 40.61
	ATOM	1131 OG SER 303 1132 C SER 305	78.597 28.664 14.589 1.00 33.02
25	ATOM		78.771 27.718 13.816 1.00 35.32
25	ATOM	1133 O SER 305 1134 N LEU 306	77.488 29.404 14.580 1.00 32.14
	ATOM	1134 N LEU 306 1135 CA LEU 306	76.391 29.121 13.653 1.00 31.02
	ATOM	1136 CB LEU 306	75.138 29.936 13.996 1.00 22.76
	ATOM	1136 CB LEU 306	74.361 29.487 15.235 1.00 24.42
20	ATOM	1137 CG LEU 306	73.094 30.311 15.380 1.00 23.13
30	ATOM	1139 CD2 LEU 306	74.016 28.009 15.126 1.00 25.53
	ATOM	1140 C LEU 306	76.780 29.354 12.198 1.00 33.11
	ATOM ATOM	1140 C LEU 300 1141 O LEU 306	76.161 28.796 11.293 1.00 32.60
		1141 O LEO 300 1142 N SER 307	77.821 30.153 11.975 1.00 36.12
25	ATOM	1142 N SER 307 1143 CA SER 307	78.296 30.448 10.624 1.00 38.80
35	ATOM	1144 CB SER 307	79.514 31.373 10.677 1.00 41.64
	ATOM	1144 CB SER 307	79.224 32.556 11.401 1.00 54.66
	ATOM		78.650 29.182 9.845 1.00 36.98
	ATOM		78.302 29.055 8.669 1.00 42.87
40	ATOM		79.315 28.239 10.509 1.00 35.72
40	ATOM	1148 N ALA 308	79.719 26.983 9.879 1.00 32.70
	ATOM	1149 CA ALA 308 1150 CB ALA 308	80.683 26.227 10.782 1.00 33.88
	ATOM		78.531 26.093 9.521 1.00 34.83
	ATOM		78.620 25.278 8.600 1.00 39.61
4 ~	ATOM	1152 O ALA 308 1153 N PHE 309	77.424 26.250 10.244 1.00 31.54
45	ATOM		76.226 25.453 9.999 1.00 32.43
	ATOM	1154 CA PHE 309	75.259 25.558 11.182 1.00 30.89
	ATOM		75.718 24.826 12.415 1.00 33.73
	ATOM		75.718 24.826 12.413 1.00 33.73 76.769 25.314 13.183 1.00 40.48
	ATOM	1157 CD1 PHE 309	75.091 23.654 12.816 1.00 35.96
50	ATOM	1158 CD2 PHE 309	/3.091 23.034 12.810 1.00 33.90

	ATOM	1159 CE1 PHE 309	77.189 24.643 14.334 1.00 37.87
	ATOM	1160 CE2 PHE 309	75.502 22.975 13.962 1.00 38.44
	ATOM	1161 CZ PHE 309	76.553 23.471 14.722 1.00 37.34
	ATOM	1162 C PHE 309	75.507 25.809 8.693 1.00 34.76
5	<b>ATOM</b>	1163 O PHE 309	74.810 24.969 8.118 1.00 36.18
	ATOM	1164 N ASN 310	75.693 27.040 8.218 1.00 35.80
	ATOM	1165 CA ASN 310	75.060 27.506 6.980 1.00 41.00
	<b>ATOM</b>	1166 CB ASN 310	75.705 26.852 5.755 1.00 51.94
	ATOM	1167 CG ASN 310	77.053 27.452 5.419 1.00 67.92
10	ATOM	1168 OD1 ASN 310	77.139 28.439 4.687 1.00 77.32
	ATOM	1169 ND2 ASN 310	78.116 26.869 5.962 1.00 72.62
	<b>ATOM</b>	1170 C ASN 310	73.560 27.245 6.985 1.00 38.15
	<b>ATOM</b>	1171 O ASN 310	73.034 26.515 6.141 1.00 35.87
	ATOM	1172 N LEU 311	72.885 27.819 7.971 1.00 33.94
15	<b>ATOM</b>	1173 CA LEU 311	71.450 27.651 8.111 1.00 32.09
	ATOM	1174 CB LEU 311	71.011 28.009 9.533 1.00 28.06
	ATOM	1175 CG LEU 311	71.656 27.301 10.724 1.00 26.38
	ATOM	1176 CD1 LEU 311	71.092 27.883 12.012 1.00 23.56
	ATOM	1177 CD2 LEU 311	71.409 25.801 10.651 1.00 21.24
20	ATOM	1178 C LEU 311	70.705 28.542 7.124 1.00 33.00
	ATOM	1179 O LEU 311	71.173 29.630 6.782 1.00 35.47
	ATOM	1180 N ASP 312	69.569 28.057 6.638 1.00 27.78
	ATOM	1181 CA ASP 312	68.749 28.841 5.733 1.00 27.06
	ATOM	1182 CB ASP 312	68.385 28.049 4.456 1.00 25.84
25	ATOM	1183 CG ASP 312	67.580 26.778 4.724 1.00 25.67
	ATOM	1184 OD1 ASP 312	67.124 26.541 5.860 1.00 28.20
	ATOM	1185 OD2 ASP 312	67.387 26.008 3.762 1.00 27.62
	ATOM	1186 C ASP 312	67.517 29.314 6.514 1.00 28.51
	ATOM	1187 O ASP 312	67.371 28.990 7.703 1.00 25.35
30	ATOM	1188 N ASP 313	66.633 30.060 5.855 1.00 22.16
	ATOM	1189 CA ASP 313	65.430 30.589 6.494 1.00 21.37
	ATOM	1190 CB ASP 313	64.625 31.431 5.499 1.00 25.11
	ATOM	1191 CG ASP 313	65.380 32.666 5.025 1.00 31.54
	ATOM	1192 OD1 ASP 313	65.119 33.115 3.890 1.00 35.35
35	ATOM	1193 OD2 ASP 313	66.225 33.193 5.783 1.00 35.37
	ATOM	1194 C ASP 313	64.524 29.535 7.120 1.00 21.11
	ATOM	1195 O ASP 313	63.904 29.783 8.158 1.00 23.68
	ATOM	1196 N THR 314	64.440 28.367 6.489 1.00 22.88
	ATOM	1197 CA THR 314	63.591 27.281 6.981 1.00 22.81
40	ATOM	1198 CB THR 314	63.472 26.155 5.927 1.00 26.00
	ATOM	1199 OG1 THR 314	62.873 26.679 4.732 1.00 20.14
	ATOM	1200 CG2 THR 314	62.629 25.010 6.457 1.00 17.51
	ATOM	1201 C THR 314	64.086 26.706 8.310 1.00 19.46
	ATOM	1202 O THR 314	63.312 26.529 9.247 1.00 19.33
45	ATOM	1203 N GLU 315	65.381 26.431 8.392 1.00 17.49
	ATOM	1204 CA GLU 315	65.965 25.885 9.611 1.00 20.62
	ATOM	1205 CB GLU 315	67.426 25.514 9.358 1.00 14.39
	ATOM	1206 CG GLU 315	67.539 24.339 8.400 1.00 13.07
	ATOM	1207 CD GLU 315	68.923 24.125 7.835 1.00 14.98
50	ATOM	1207 CB GEC 313	69.634 25.116 7.552 1.00 17.71

	ATOM	1209 OE2 GLU 315	69.287 22.948 7.651 1.00 17.88
	ATOM	1210 C GLU 315	65.810 26.883 10.762 1.00 20.57
	ATOM	1211 O GLU 315	65.368 26.518 11.854 1.00 18.43
	ATOM	1212 N VAL 316	66.096 28.154 10.488 1.00 19.19
5	ATOM	1213 CA VAL 316	65.955 29.203 11.490 1.00 16.53
	ATOM	1214 CB VAL 316	66.418 30.567 10.933 1.00 17.42
	ATOM	1215 CG1 VAL 316	66.149 31.687 11.940 1.00 13.89
	ATOM	1216 CG2 VAL 316	67.900 30.506 10.594 1.00 14.31
	ATOM	1217 C VAL 316	64.488 29.291 11.927 1.00 19.53
10	<b>ATOM</b>	1218 O VAL 316	64.191 29.448 13.110 1.00 19.86
	<b>ATOM</b>	1219 N ALA 317	63.575 29.159 10.970 1.00 19.02
	ATOM	1220 CA ALA 317	62.145 29.215 11.254 1.00 16.95
	ATOM	1221 CB ALA 317	61.357 29.239 9.951 1.00 17.68
	<b>ATOM</b>	1222 C ALA 317	61.674 28.047 12.126 1.00 14.13
15	<b>ATOM</b>	1223 O ALA 317	60.875 28.228 13.045 1.00 15.34
	<b>ATOM</b>	1224 N LEU 318	62.154 26.847 11.819 1.00 17.41
	ATOM	1225 CA LEU 318	61.769 25.653 12.569 1.00 19.10
	ATOM	1226 CB LEU 318	62.186 24.398 11.802 1.00 18.21
	ATOM	1227 CG LEU 318	61.443 24.209 10.473 1.00 19.02
20	ATOM	1228 CD1 LEU 318	62.105 23.128 9.646 1.00 16.10
	ATOM	1229 CD2 LEU 318	59.987 23.875 10.735 1.00 11.32
	ATOM	1230 C LEU 318	62.399 25.685 13.954 1.00 22.38
	ATOM	1231 O LEU 318	61.782 25.278 14.945 1.00 21.64
	ATOM	1232 N LEU 319	63.619 26.207 14.016 1.00 20.97
25	ATOM	1233 CA LEU 319	64.338 26.344 15.270 1.00 19.71
	ATOM	1234 CB LEU 319	65.715 26.951 15.005 1.00 20.56
	ATOM	1235 CG LEU 319	66.722 27.036 16.152 1.00 32.05
	ATOM	1236 CD1 LEU 319	66.704 25.760 16.963 1.00 33.15
	ATOM	1237 CD2 LEU 319	68.109 27.303 15.590 1.00 28.25
30	ATOM	1238 C LEU 319	63.496 27.254 16.164 1.00 20.66
	ATOM	1239 O LEU 319	63.215 26.920 17.313 1.00 24.47
	ATOM	1240 N GLN 320	63.026 28.365 15.604 1.00 19.25
	ATOM	1241 CA GLN 320	62.191 29.307 16.346 1.00 19.02
26	ATOM	1242 CB GLN 320	61.842 30.526 15.488 1.00 19.11 63.032 31.377 15.101 1.00 20.02
35	ATOM	1243 CG GLN 320	63.032 31.377 15.101 1.00 20.02 62.665 32.562 14.224 1.00 23.65
	ATOM ATOM	1244 CD GLN 320 1245 OEI GLN 320	62.663 32.362 14.224 1.00 23.03
	ATOM	1246 NE2 GLN 320	61.440 32.574 13.704 1.00 20.77
	ATOM	1246 NE2 GLN 320 1247 C GLN 320	60.905 28.635 16.811 1.00 20.52
40	ATOM	1247 C GLN 320 1248 O GLN 320	60.465 28.845 17.938 1.00 22.04
40	ATOM	1249 N ALA 321	60.306 27.825 15.942 1.00 21.01
	ATOM	1250 CA ALA 321	59.069 27.128 16.280 1.00 16.83
	ATOM	1250 CA ALA 321 1251 CB ALA 321	58.556 26.358 15.079 1.00 16.58
	ATOM	1251 CB ALA 321	59.288 26.185 17.462 1.00 18.15
45	ATOM	1252 C ALA 321	58.427 26.069 18.344 1.00 13.03
	ATOM	1254 N VAL 322	60.442 25.523 17.481 1.00 14.89
	ATOM	1255 CA VAL 322	60.774 24.599 18.559 1.00 19.05
	ATOM	1256 CB VAL 322	62.051 23.779 18.233 1.00 21.50
	ATOM	1257 CG1 VAL 322	62.510 22.990 19.457 1.00 21.49
50	ATOM	1258 CG2 VAL 322	61.773 22.819 17.073 1.00 15.42

	ATOM	1259 C VAL 322	60.947 25.375 19.867 1.00 19.89
	ATOM	1260 O VAL 322	60.478 24.940 20.919 1.00 21.58
	ATOM	1261 N LEU 323	61.591 26.537 19.788 1.00 20.25
	ATOM	1262 CA LEU 323	61.804 27.387 20.959 1.00 19.32
5	ATOM	1263 CB LEU 323	62.683 28.586 20.597 1.00 12.95
3		1264 CG LEU 323	64.129 28.273 20.217 1.00 20.70
	ATOM	1265 CD1 LEU 323	64.805 29.503 19.641 1.00 13.23
	ATOM		64.883 27.767 21.438 1.00 22.91
	ATOM		60.468 27.884 21.497 1.00 20.25
10	ATOM	1267 C LEU 323	
10	ATOM	1268 O LEU 323	
	ATOM	1269 N LEU 324	
	ATOM	1270 CA LEU 324	
	ATOM	1271 CB LEU 324	57.555 29.333 19.707 1.00 18.45
	ATOM	1272 CG LEU 324	56.119 29.847 19.868 1.00 17.07
15	ATOM	1273 CD1 LEU 324	56.083 31.092 20.752 1.00 15.39
	ATOM	1274 CD2 LEU 324	55.545 30.162 18.498 1.00 17.90
	ATOM	1275 C LEU 324	57.342 27.706 21.598 1.00 21.54
	ATOM	1276 O LEU 324	56.742 27.967 22.642 1.00 23.41
	ATOM	1277 N MET 325	57.249 26.521 21.003 1.00 24.63
20	ATOM	1278 CA MET 325	56.380 25.476 21.545 1.00 25.35
	ATOM	1279 CB MET 325	55.901 24.536 20.430 1.00 25.53
	ATOM	1280 CG MET 325	55.235 25.220 19.232 1.00 21.89
	ATOM	1281 SD MET 325	53.871 26.337 19.649 1.00 25.50
	ATOM	1282 CE MET 325	52.705 25.250 20.397 1.00 17.66
25	ATOM	1283 C MET 325	57.031 24.676 22.675 1.00 27.58
	ATOM	1284 O MET 325	56.988 23.450 22.690 1.00 28.61
	ATOM	1285 N SER 326	57.613 25.376 23.638 1.00 27.98
	ATOM	1286 CA SER 326	58.265 24.718 24.757 1.00 31.60
	ATOM	1287 CB SER 326	59.527 25.493 25.155 1.00 35.80
30	ATOM	1288 OG SER 326	60.123 24.966 26.327 1.00 43.74
	ATOM	1289 C SER 326	57.313 24.624 25.939 1.00 32.12
	ATOM	1290 O SER 326	56.590 25.574 26.240 1.00 30.91
	ATOM	1291 N THR 327	57.276 23.464 26.583 1.00 35.41
	ATOM	1292 CA THR 327	56.420 23.278 27.747 1.00 39.61
35	ATOM	1293 CB THR 327	55.777 21.890 27.758 1.00 38.84
	ATOM	1294 OG1 THR 327	56.784 20.890 27.538 1.00 42.53
	ATOM	1295 CG2 THR 327	54.716 21.802 26.679 1.00 40.78
	ATOM	1296 C THR 327	57.232 23.471 29.022 1.00 43.86
	ATOM	1297 O THR 327	56.785 23.133 30.118 1.00 42.40
40	ATOM	1298 N ASP 328	58.417 24.054 28.869 1.00 47.35
	ATOM	1299 CA ASP 328	59.309 24.308 29.987 1.00 49.43
	ATOM	1300 CB ASP 328	60.750 24.358 29.482 1.00 58.03
	ATOM	1301 CG ASP 328	61.718 23.687 30.425 1.00 72.16
	ATOM	1302 OD1 ASP 328	61.816 24.117 31.595 1.00 82.32
45	ATOM	1303 OD2 ASP 328	62.378 22.720 29.994 1.00 81.63
	ATOM	1304 C ASP 328	58.951 25.625 30.676 1.00 47.99
	ATOM	1305 O ASP 328	59.830 26.373 31.093 1.00 53.33
	ATOM	1306 N ARG 329	57.657 25.910 30.780 1.00 48.33
	ATOM	1307 CA ARG 329	57.177 27.135 31.413 1.00 47.67
50	ATOM	1308 CB ARG 329	56.562 28.091 30.379 1.00 47.64

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ATOM 1309 CG ARG 329
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     ATOM 1310 CD ARG 329
     ATOM 1311 NE ARG 329
                                58.759 28.682 27.288 1.00 41.17
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     ATOM 1313 NH1 ARG 329
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     ATOM 1329 CG LEU 332
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     ATOM 1331 CD2 LEU 332
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     ATOM 1333 O LEU 332
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     ATOM 1334 N LEU 333
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     ATOM 1337 CG LEU 333
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     ATOM 1352 CG1 VAL 335
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     ATOM 1358 CB ASP 336
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PCT/US98/25296

### WO 99/26966

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     ATOM 1381 CD GLU 339
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     ATOM 1408 CB GLU 343
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PCT/US98/25296

WO 99/26966

ATOM 1409 C GLU 343 57.225 17.094 17.119 1.00 19.18 ATOM 1410 O GLU 343 58.156 16.743 16.393 1.00 21.11 56.077 17.570 16.648 1.00 19.93 ATOM 1411 N ALA 344 55.803 17.662 15.217 1.00 20.20 ATOM 1412 CA ALA 344 54.411 18.216 14.989 1.00 16.46 ATOM 1413 CB ALA 344 56.850 18.539 14.528 1.00 20.75 ATOM 1414 C ALA 344 57.432 18.140 13.514 1.00 25.13 ATOM 1415 O ALA 344 ATOM 1416 N TYR 345 57.105 19.722 15.088 1.00 21.31 ATOM 1417 CA TYR 345 58.107 20.631 14.531 1.00 15.93 58.127 21.969 15.282 1.00 17.29 ATOM 1418 CB TYR 345 10 ATOM 1419 CG TYR 345 57.049 22.927 14.833 1.00 16.11 56.017 23.296 15.689 1.00 9.93 ATOM 1420 CD1 TYR 345 ATOM 1421 CE1 TYR 345 54.999 24.138 15.263 1.00 16.95 ATOM 1422 CD2 TYR 345 57.041 23.431 13.531 1.00 19.84 ATOM 1423 CE2 TYR 345 56.026 24.276 13.094 1.00 17.13 15 55.005 24.622 13.963 1.00 18.12 ATOM 1424 CZ TYR 345 53.980 25.430 13.530 1.00 26.25 ATOM 1425 OH TYR 345 59.493 20.008 14.554 1.00 20.65 ATOM 1426 C TYR 345 ATOM 1427 O TYR 345 60.240 20.129 13.583 1.00 20.75 20 ATOM 1428 N LEU 346 59.832 19.337 15.655 1.00 22.14 ATOM 1429 CA LEU 346 61.134 18.684 15.803 1.00 19.43 61.267 18.041 17.186 1.00 19.92 ATOM 1430 CB LEU 346 ATOM 1431 CG LEU 346 61.683 18.945 18.347 1.00 25.56 ATOM 1432 CD1 LEU 346 61.440 18.244 19.677 1.00 22.06 63.147 19.332 18.197 1.00 17.62 ATOM 1433 CD2 LEU 346 25 ATOM 1434 C LEU 346 61.359 17.635 14.723 1.00 19.30 62.441 17.560 14.142 1.00 22.84 ATOM 1435 O LEU 346 ATOM 1436 N LEU 347 60.337 16.826 14.456 1.00 25.17 ATOM 1437 CA LEU 347 60,423 15,790 13,427 1.00 24,55 ATOM 1438 CB LEU 347 59.187 14.892 13.453 1.00 25.47 30 59.256 13.654 14.345 1.00 30.65 ATOM 1439 CG LEU 347 ATOM 1440 CD1 LEU 347 57.941 12.890 14.258 1.00 34.28 60.416 12.765 13.908 1.00 28.26 ATOM 1441 CD2 LEU 347 60.584 16.400 12.042 1.00 24.00 ATOM 1442 C LEU 347 ATOM 1443 O LEU 347 61.399 15.932 11.245 1.00 29.74 35 59.809 17.443 11.761 1.00 22.72 ATOM 1444 N ALA 348 59.875 18.125 10.475 1.00 19.19 ATOM 1445 CA ALA 348 58.789 19.188 10.388 1.00 22.73 ATOM 1446 CB ALA 348 61,246 18,762 10,316 1,00 20,34 ATOM 1447 C ALA 348 ATOM 1448 O ALA 348 61.881 18.633 9.274 1.00 23.94 40 61.707 19.402 11.388 1.00 22.19 ATOM 1449 N PHE 349 ATOM 1450 CA PHE 349 63.001 20.078 11.435 1.00 19.41 63.185 20.701 12.832 1.00 17.45 ATOM 1451 CB PHE 349 64.371 21.632 12.963 1.00 18.70 ATOM 1452 CG PHE 349 65.183 21.943 11.874 1.00 19.09 ATOM 1453 CD1 PHE 349 64.669 22.203 14.199 1.00 21.81 ATOM 1454 CD2 PHE 349 66.270 22.811 12.012 1.00 21.49 ATOM 1455 CE1 PHE 349 ATOM 1456 CE2 PHE 349 65.753 23.072 14.351 1.00 18.58 66.555 23.376 13.256 1.00 18.67 ATOM 1457 CZ PHE 349 64.110 19.071 11.136 1.00 20.96 50 ATOM 1458 C PHE 349

			C4 0C7 10 211 10 202 1 00 25 10
	ATOM	1459 O PHE 349	64.967 19.311 10.283 1.00 25.19
	ATOM	1460 N GLU 350	64.076 17.935 11.824 1.00 23.96
	ATOM	1461 CA GLU 350	65.077 16.888 11.642 1.00 27.98
_	ATOM	1462 CB GLU 350	64.794 15.721 12.591 1.00 28.90 65.738 14.542 12.413 1.00 39.36
5	ATOM	1463 CG GLU 350	001,00 1 110 12 12111
	ATOM	1464 CD GLU 350	65.603 13.497 13.505 1.00 41.62
	ATOM	1465 OE1 GLU 350	64.475 13.260 13.988 1.00 43.67
	ATOM	1466 OE2 GLU 350	66.636 12.908 13.876 1.00 49.64
	ATOM	1467 C GLU 350	65.100 16.385 10.203 1.00 27.12
10	ATOM	1468 O GLU 350	66.158 16.288 9.577 1.00 27.44
	ATOM	1469 N HIS 351	63.918 16.088 9.678 1.00 27.36
	ATOM	1470 CA HIS 351	63.787 15.591 8.318 1.00 23.97
	ATOM	1471 CB HIS 351	62.366 15.087 8.090 1.00 22.89
	ATOM	1472 CG HIS 351	61.991 13.945 8.986 1.00 24.58
15	ATOM	1473 CD2 HIS 351	62.736 13.209 9.846 1.00 25.83
	ATOM	1474 ND1 HIS 351	60.709 13.448 9.073 1.00 26.50
	ATOM	1475 CEI HIS 351	60.677 12.460 9.948 1.00 24.81
	ATOM	1476 NE2 HIS 351	61.896 12.295 10.431 1.00 28.42
	ATOM	1477 C HIS 351	64.200 16.635 7.278 1.00 24.22
20	ATOM	1478 O HIS 351	64.757 16.287 6.236 1.00 25.79
	ATOM	1479 N TYR 352	63.969 17.912 7.572 1.00 21.04
	ATOM	1480 CA TYR 352	64.363 18.974 6.654 1.00 18.98
	ATOM	1481 CB TYR 352	63.770 20.321 7.067 1.00 17.08
	ATOM	1482 CG TYR 352	64.127 21.413 6.090 1.00 21.83
25	ATOM	1483 CD1 TYR 352	63.537 21.467 4.828 1.00 20.07
	ATOM	1484 CE1 TYR 352	63.941 22.411 3.883 1.00 23.51
	ATOM	1485 CD2 TYR 352	65.121 22.339 6.388 1.00 19.94
	ATOM	1486 CE2 TYR 352	65.531 23.284 5.452 1.00 20.85
	ATOM	1487 CZ TYR 352	64.942 23.313 4.203 1.00 24.80
30	ATOM	1488 OH TYR 352	65.380 24.221 3.269 1.00 26.74
	ATOM	1489 C TYR 352	65.889 19.055 6.624 1.00 20.58
	ATOM	1490 O TYR 352	66.492 19.276 5.570 1.00 22.72
	ATOM	1491 N VAL 353	66.508 18.877 7.789 1.00 28.34
	ATOM	1492 CA VAL 353	67.967 18.892 7.904 1.00 22.38
35	ATOM	1493 CB VAL 353	68.419 18.755 9.389 1.00 26.46
	ATOM	1494 CG1 VAL 353	69.915 18.527 9.478 1.00 20.92
	ATOM	1495 CG2 VAL 353	68.053 20.009 10.165 1.00 22.46
	ATOM	1496 C VAL 353	68.518 17.725 7.078 1.00 23.51
	<b>ATOM</b>	1497 O VAL 353	69.535 17.865 6.391 1.00 24.73
40	<b>ATOM</b>	1498 N ASN 354	67.850 16.575 7.158 1.00 20.93
	ATOM	1499 CA ASN 354	68.252 15.392 6.397 1.00 27.25
	ATOM	1500 CB ASN 354	67.320 14.210 6.680 1.00 28.43
	ATOM	1501 CG ASN 354	67.521 13.607 8.058 1.00 31.50
	ATOM	1502 OD1 ASN 354	
45	ATOM	1503 ND2 ASN 354	
	ATOM	1504 C ASN 354	68.182 15.721 4.908 1.00 31.27
	ATOM	1505 O ASN 354	69.066 15.347 4.134 1.00 34.22
	ATOM	1506 N HIS 355	67.124 16.429 4.520 1.00 30.49
	ATOM	1507 CA HIS 355	66.917 16.826 3.132 1.00 26.88
50	ATOM	1508 CB HIS 355	65.548 17.494 2.975 1.00 27.27

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65.319 18.103 1.625 1.00 37.76
    ATOM 1509 CG HIS 355
    ATOM 1510 CD2 HIS 355
                               65.439 19.382 1.196 1.00 35.28
                               64.913 17.369 0.532 1.00 34.93
    ATOM 1511 ND1 HIS 355
                               64.789 18.169 -0.513 1.00 34.84
    ATOM 1512 CE1 HIS 355
                               65.104 19.394 -0.135 1.00 33.13
    ATOM 1513 NE2 HIS 355
                              68.016 17.748 2.610 1.00 24.66
    ATOM 1514 C HIS 355
                              68.420 17.630 1.456 1.00 26.62
    ATOM 1515 O HIS 355
                               68.487 18.670 3.448 1.00 25.86
    ATOM 1516 N ARG 356
                               69.536 19.608 3.040 1.00 26.94
    ATOM 1517 CA ARG 356
                                69.620 20.791 3.996 1.00 20.57
    ATOM 1518 CB ARG 356
10
                                68.453 21.727 3.899 1.00 19.69
    ATOM 1519 CG ARG 356
                                68.866 23.110 4.340 1.00 23.81
    ATOM 1520 CD ARG 356
                                69.768 23.746 3.388 1.00 23.14
    ATOM 1521 NE ARG 356
    ATOM 1522 CZ ARG 356
                                70.641 24.697 3.702 1.00 24.11
    ATOM 1523 NH1 ARG 356
                                70.755 25.129 4.949 1.00 26.29
15
                                71.384 25.242 2.754 1.00 32.79
    ATOM 1524 NH2 ARG 356
                               70.921 19.002 2.875 1.00 29.38
    ATOM 1525 C ARG 356
                               71.795 19.607 2.257 1.00 32.91
    ATOM 1526 O ARG 356
                               71.133 17.848 3.498 1.00 33.39
    ATOM 1527 N LYS 357
                                72.401 17.128 3.417 1.00 35.97
    ATOM 1528 CA LYS 357
20
                               72.479 16.363 2.089 1.00 40.55
    ATOM 1529 CB LYS 357
                                71.327 15.381 1.891 1.00 44.03
    ATOM 1530 CG LYS 357
                                71.360 14.722 0.523 1.00 52.31
    ATOM 1531 CD LYS 357
                               70.171 13.787 0.343 1.00 56.99
    ATOM 1532 CE LYS 357
                               70.208 13.085 -0.970 1.00 64.78
     ATOM 1533 NZ LYS 357
25
                               73.657 17.981 3.629 1.00 38.55
    ATOM 1534 C LYS 357
                               74.518 18.079 2.748 1.00 42.50
     ATOM 1535 O LYS 357
                              73.751 18.601 4.802 1.00 35.00
     ATOM 1536 N HIS 358
     ATOM 1537 CA HIS 358
                               74.906 19.418 5.155 1.00 32.94
     ATOM 1538 CB HIS 358
                               74,732 20.018 6.552 1.00 27.62
30
     ATOM 1539 CG HIS 358
                               73.669 21.067 6.643 1.00 26.64
                                72.330 20.968 6.819 1.00 20.85
     ATOM 1540 CD2 HIS 358
                                73.950 22.416 6.587 1.00 24.71
     ATOM 1541 ND1 HIS 358
                               72.831 23.103 6.724 1.00 21.02
     ATOM 1542 CE1 HIS 358
                               71.834 22.248 6.865 1.00 21.42
     ATOM 1543 NE2 HIS 358
35
                              76.140 18.520 5.176 1.00 36.60
     ATOM 1544 C HIS 358
                               76.072 17.379 5.635 1.00 38.73
     ATOM 1545 O HIS 358
                               77.267 19.037 4.702 1.00 41.40
     ATOM 1546 N ASN 359
                                78.515 18.277 4.689 1.00 45.02
     ATOM 1547 CA ASN 359
                                79.441 18.799 3.587 1.00 42.57
     ATOM 1548 CB ASN 359
40
     ATOM 1549 C ASN 359
                               79.193 18.386 6.058 1.00 46.59
     ATOM 1550 O ASN 359
                               80.405 18.588 6.150 1.00 52.31
                               78.400 18.254 7.117 1.00 45.14
     ATOM 1551 N ILE 360
                               78.896 18.348 8.487 1.00 43.69
     ATOM 1552 CA ILE 360
                               78.330 19.597 9.207 1.00 40.08
     ATOM 1553 CB ILE 360
45
                                78.824 19.657 10.645 1.00 32.11
     ATOM 1554 CG2 ILE 360
                                78.733 20.864 8.452 1.00 41.47
     ATOM 1555 CG1 ILE 360
                                78.057 22.115 8.954 1.00 44.93
     ATOM 1556 CD1 ILE 360
                              78.452 17.101 9.242 1.00 43.63
     ATOM 1557 C ILE 360
                              77.257 16.797 9.313 1.00 45.20
     ATOM 1558 O ILE 360
50
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ATOM 1559 N PRO 361
                               79.413 16.337 9.780 1.00 43.91
     ATOM 1560 CD PRO 361
                                80.871 16.540 9.699 1.00 47.07
     ATOM 1561 CA PRO 361
                                79.087 15.118 10.526 1.00 41.66
                                80.462 14.495 10.782 1.00 43.73
     ATOM 1562 CB PRO 361
     ATOM 1563 CG PRO 361
                                81.383 15.679 10.830 1.00 45.45
     ATOM 1564 C PRO 361
                               78.332 15.403 11.832 1.00 36.42
     ATOM 1565 O PRO 361
                               78.679 16.325 12.572 1.00 35.74
     ATOM 1566 N HIS 362
                               77.291 14.610 12.088 1.00 33.14
     ATOM 1567 CA HIS 362
                               76.462 14.726 13.292 1.00 34.09
                                77.288 14.413 14.547 1.00 33.82
     ATOM 1568 CB HIS 362
10
                                78.132 13.181 14.424 1.00 36.04
     ATOM 1569 CG HIS 362
                                77.793 11.885 14.224 1.00 34.77
     ATOM 1570 CD2 HIS 362
                                79.509 13.212 14.482 1.00 37.16
     ATOM 1571 ND1 HIS 362
     ATOM 1572 CE1 HIS 362
                                79.983 11.990 14.325 1.00 37.16
                                78.962 11.165 14.167 1.00 40.13
15
     ATOM 1573 NE2 HIS 362
     ATOM 1574 C HIS 362
                               75.829 16.110 13.417 1.00 31.00
     ATOM 1575 O HIS 362
                               75.617 16.608 14.525 1.00 30.22
     ATOM 1576 N PHE 363
                               75.478 16.690 12.272 1.00 33.06
     ATOM 1577 CA PHE 363
                                74.878 18.021 12.200 1.00 28.08
                                74.503 18.355 10.747 1.00 25.26
20
     ATOM 1578 CB PHE 363
     ATOM 1579 CG PHE 363
                                73.923 19.733 10.567 1.00 24.91
                                74.750 20.817 10.320 1.00 27.60
     ATOM 1580 CD1 PHE 363
     ATOM 1581 CD2 PHE 363
                                72.552 19.948 10.664 1.00 25.52
                                74.221 22.100 10.175 1.00 29.70
     ATOM 1582 CE1 PHE 363
     ATOM 1583 CE2 PHE 363
                                72.014 21.227 10.522 1.00 25.88
25
     ATOM 1584 CZ PHE 363
                                72.850 22.304 10.278 1.00 21.49
     ATOM 1585 C PHE 363
                               73.659 18.201 13.099 1.00 23.79
     ATOM 1586 O PHE 363
                               73.587 19.164 13.863 1.00 24.48
     ATOM 1587 N TRP 364
                               72.707 17.277 13.012 1.00 23.13
     ATOM 1588 CA TRP 364
                                71.484 17.369 13.805 1.00 25.06
30
                                70.536 16.201 13.494 1.00 21.17
     ATOM 1589 CB TRP
                          364
     ATOM 1590 CG TRP 364
                                69.247 16.220 14.271 1.00 23.14
     ATOM 1591 CD2 TRP 364
                                68.261 17.266 14.296 1.00 27.68
     ATOM 1592 CE2 TRP 364
                                67.229 16.845 15.165 1.00 28.31
     ATOM 1593 CE3 TRP
                          364
                                68.149 18.517 13.671 1.00 26.46
                                68.784 15.241 15.096 1.00 23.76
     ATOM 1594 CD1 TRP 364
     ATOM 1595 NE1 TRP 364
                                67.576 15.607 15.637 1.00 32.12
     ATOM 1596 CZ2 TRP 364
                                66.100 17.628 15.427 1.00 25.63
     ATOM 1597 CZ3 TRP 364
                                67.028 19.294 13.931 1.00 25.55
40 - ATOM 1598 CH2 TRP 364
                                66.017 18.845 14.803 1.00 29.79
     ATOM 1599 C TRP 364
                               71.715 17.531 15.312 1.00 27.80
     ATOM 1600 O TRP 364
                               71.212 18.486 15.904 1.00 26.96
     ATOM 1601 N PRO 365
                               72.458 16.605 15.955 1.00 30.69
     ATOM 1602 CD PRO 365
                                72.974 15.308 15.481 1.00 31.45
45
     ATOM 1603 CA PRO 365
                                72.687 16.757 17.397 1.00 27.97
     ATOM 1604 CB PRO 365
                                73.506 15.512 17.752 1.00 26.50
     ATOM 1605 CG PRO 365
                                73.057 14.509 16.757 1.00 33.47
     ATOM 1606 C PRO 365
                               73.457 18.043 17.709 1.00 27.10
     ATOM 1607 O PRO 365
                               73.154 18.736 18.681 1.00 26.88
     ATOM 1608 N LYS 366
50
                               74.440 18.365 16.873 1.00 26.99
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	A TO 3.4	1600 CA TVC 366	75.230 19.577 17.061 1.00 30.69
	ATOM ATOM	1609 CA LYS 366 1610 CB LYS 366	76.275 19.708 15.957 1.00 28.53
	ATOM	1611 CG LYS 366	77.481 18.804 16.106 1.00 28.89
	ATOM	1612 CD LYS 366	78.430 19.027 14.939 1.00 32.51
_	ATOM	1612 CD L13 366	79.743 18.294 15.116 1.00 38.52
5		1614 NZ LYS 366	80,632 18.506 13.939 1.00 45.28
	ATOM		74.349 20.831 17.079 1.00 36.18
	ATOM		74.472 21.672 17.972 1.00 39.82
	ATOM	1616 O LYS 366	73.464 20.950 16.091 1.00 37.54
10	ATOM	1617 N LEU 367	72.557 22.092 15.994 1.00 36.14
10	ATOM	1618 CA LEU 367	71.803 22.070 14.659 1.00 32.20
	ATOM	1619 CB LEU 367	
	ATOM	1620 CG LEU 367	70.764 23.179 14.447 1.00 36.16
	ATOM	1621 CD1 LEU 367	71.402 24.567 14.618 1.00 20.60
	ATOM	1622 CD2 LEU 367	70.139 23.030 13.065 1.00 34.30
15	ATOM	1623 C LEU 367	71.561 22.060 17.143 1.00 36.84
	ATOM	1624 O LEU 367	71.231 23.091 17.729 1.00 36.94
	ATOM	1625 N LEU 368	71.083 20.866 17.459 1.00 37.81
	ATOM	1626 CA LEU 368	70.130 20.683 18.536 1.00 34.83
	ATOM	1627 CB LEU 368	69.763 19.205 18.622 1.00 36.98
20	ATOM	1628 CG LEU 368	68.421 18.777 19.205 1.00 40.34
	ATOM	1629 CD1 LEU 368	67.276 19.595 18.619 1.00 36.28
	ATOM	1630 CD2 LEU 368	68.241 17.299 18.908 1.00 39.39
	ATOM	1631 C LEU 368	70.755 21.182 19.843 1.00 38.32
	ATOM	1632 O LEU 368	70.059 21.711 20.707 1.00 41.87
25	ATOM	1633 N MET 369	72.075 21.057 19.962 1.00 39.46
	ATOM	1634 CA MET 369	72.790 21.515 21.154 1.00 40.12
	ATOM	1635 CB MET 369	74.219 20.971 21.168 1.00 41.26
	ATOM	1636 CG MET 369	74.307 19.493 21.521 1.00 47.83
	ATOM	1637 SD MET 369	75.961 18.810 21.289 1.00 55.72
30	ATOM	1638 CE MET 369	76.809 19.474 22.727 1.00 54.37
	ATOM	1639 C MET 369	72.805 23.039 21.251 1.00 42.81
	ATOM	1640 O MET 369	72.990 23.601 22.335 1.00 47.81
	ATOM	1641 N LYS 370	72.622 23.708 20.115 1.00 40.09
	ATOM	1642 CA LYS 370	72.588 25.165 20.080 1.00 33.65
35	ATOM	1643 CB LYS 370	72.751 25.677 18.650 1.00 30.83
	ATOM	1644 CG LYS 370	74.138 25.435 18.078 1.00 30.98
	ATOM	1645 CD LYS 370	75.188 26.198 18.867 1.00 37.82
	ATOM	1646 CE LYS 370	76.591 25.938 18.351 1.00 36.05
	ATOM	1647 NZ LYS 370	77.034 24.562 18.667 1.00 48.68
40	ATOM	1648 C LYS 370	71.293 25.684 20.702 1.00 33.32
	ATOM	1649 O LYS 370	71.218 26.842 21.112 1.00 34.75
	ATOM	1650 N VAL 371	70.277 24.826 20.779 1.00 31.90
	ATOM	1651 CA VAL 371	69.006 25.197 21.395 1.00 31.77
	ATOM	1652 CB VAL 371	67.933 24.092 21.214 1.00 30.28
45	ATOM	1653 CG1 VAL 371	66.673 24.429 21.995 1.00 30.02
	ATOM	1654 CG2 VAL 371	67.596 23.933 19.746 1.00 32.23
	ATOM	1655 C VAL 371	69.277 25.417 22.885 1.00 34.44
	ATOM	1656 O VAL 371	68.722 26.331 23.499 1.00 33.35
	ATOM	1657 N THR 372	70.161 24.590 23.443 1.00 33.15
50	ATOM	1658 CA THR 372	70.551 24.675 24.847 1.00 32.47

	ATOM	1659 CB THR 372	71.541 23.556 25.207 1.00 32.11
	ATOM	1660 OG1 THR 372	70.955 22.288 24.891 1.00 35.33
	ATOM	1661 CG2 THR 372	71.894 23.603 26.688 1.00 32.54
	ATOM	1662 C THR 372	71.226 26.020 25.108 1.00 34.49
5	ATOM	1663 O THR 372	70.936 26.696 26.099 1.00 34.07
5	ATOM	1664 N ASP 373	72.120 26.405 24.202 1.00 32.77
	ATOM	1665 CA ASP 373	72.830 27.671 24.315 1.00 28.08
	ATOM	1666 CB ASP 373	73.803 27.841 23.147 1.00 31.59
	ATOM	1667 CG ASP 373	74.910 26.789 23.142 1.00 37.29
10	ATOM	1668 OD1 ASP 373	75.170 26.169 24.196 1.00 40.82
10		1669 OD2 ASP 373	75.531 26.586 22.079 1.00 40.81
	ATOM	1670 C ASP 373	71.830 28.821 24.353 1.00 29.21
	ATOM		71.830 28.821 24.333 1.00 29.21 71.931 29.709 25.200 1.00 31.85
	ATOM		70.843 28.775 23.463 1.00 24.71
1.5	ATOM		69.813 29.802 23.403 1.00 25.25
15	ATOM		68.906 29.587 22.188 1.00 25.61
	ATOM	1674 CB LEU 374	69.480 30.084 20.858 1.00 25.51
	ATOM	1675 CG LEU 374	68.741 29.469 19.677 1.00 23.53
	ATOM	1676 CD1 LEU 374	68.741 29.469 19.677 1.00 23.33 69.405 31.596 20.820 1.00 21.92
•	ATOM	1677 CD2 LEU 374	68.994 29.827 24.686 1.00 26.84
20	ATOM	1678 C LEU 374	
	ATOM	1679 O LEU 374	68.591 30.895 25.151 1.00 28.96 68.746 28.651 25.254 1.00 31.00
	ATOM	1680 N ARG 375	67.996 28.554 26.502 1.00 32.86
	ATOM	1681 CA ARG 375	
	ATOM	1682 CB ARG 375	67.831 27.090 26.924 1.00 36.80 66.861 26.297 26.071 1.00 44.91
25	ATOM	1683 CG ARG 375	
	ATOM	1684 CD ARG 375	65.433 26.731 26.338 1.00 58.99
	ATOM	1685 NE ARG 375	64.501 26.210 25.342 1.00 72.26
	ATOM	1686 CZ ARG 375	63.909 25.020 25.404 1.00 77.46
• •	ATOM	1687 NH1 ARG 375	
30	ATOM	1688 NH2 ARG 375	
	ATOM	1689 C ARG 375	68.771 29.317 27.570 1.00 32.27
	ATOM	1690 O ARG 375	68.199 30.125 28.304 1.00 33.75
	ATOM	1691 N MET 376	70.084 29.098 27.602 1.00 32.65
	ATOM	1692 CA MET 376	70.967 29.753 28.560 1.00 35.83
35	ATOM	1693 CB MET 376	72.392 29.210 28.434 1.00 39.25
	ATOM	1694 CG MET 376	72.526 27.751 28.839 1.00 54.45
	ATOM	1695 SD MET 376	74.245 27.212 28.944 1.00 73.93
	ATOM	1696 CE MET 376	74.421 26.270 27.434 1.00 67.01
	ATOM	1697 C MET 376	70.960 31.267 28.378 1.00 35.38
40	ATOM	1698 O MET 376	70.882 32.015 29.353 1.00 34.73
	ATOM	1699 N ILE 377	71.038 31.716 27.129 1.00 32.51
	ATOM	1700 CA ILE 377	71.016 33.142 26.816 1.00 26.55
	ATOM	1701 CB ILE 377	71.182 33.370 25.299 1.00 24.84
	ATOM	1702 CG2 ILE 377	70.817 34.797 24.923 1.00 26.63
45	ATOM	1703 CG1 ILE 377	72.616 33.038 24.890 1.00 20.66
	ATOM	1704 CD1 ILE 377	72.872 33.104 23.409 1.00 20.74
	ATOM	1705 C ILE 377	69.706 33.755 27.313 1.00 25.47
	ATOM	1706 O ILE 377	69.696 34.848 27.881 1.00 29.99
	ATOM	1707 N GLY 378	68.608 33.033 27.127 1.00 25.11
50	ATOM	1708 CA GLY 378	67.321 33.522 27.580 1.00 27.82

	ATOM		7.279 33.613 29.095 1.00 30.90 6.749 34.579 29.651 1.00 31.19
	ATOM ATOM		7.851 32.611 29.761 1.00 31.62
			67.896 32.547 31.223 1.00 30.74
_	ATOM		68.433 31.198 31.671 1.00 30.82
5	ATOM		8.756 33.668 31.801 1.00 30.07
	ATOM		
	ATOM		
	ATOM		9.966 33.817 31.273 1.00 29.72 70.873 34.866 31.723 1.00 33.36
	ATOM		
10	ATOM		72.201 34.809 30.963 1.00 38.31
	ATOM		73.249 33.407 31.386 1.00 50.99
	ATOM		74.982 33.655 29.929 1.00 70.37
	ATOM		0.226 36.232 31.535 1.00 33.40
	ATOM		0.246 37.062 32.442 1.00 36.41
15	ATOM		2.615 36.456 30.374 1.00 32.55
	ATOM		8.965 37.734 30.114 1.00 26.41
	ATOM		8.434 37.811 28.681 1.00 20.89
	ATOM		7.593 39.023 28.423 1.00 15.78
• •	ATOM		57.928 40.277 28.041 1.00 12.67
20	ATOM		66.226 39.031 28.605 1.00 17.88
	ATOM		55.756 40.239 28.353 1.00 16.27
	ATOM		56.768 41.013 28.008 1.00 17.18
	ATOM		.839 38.023 31.102 1.00 26.73
0.5	ATOM		7.621 39.176 31.464 1.00 30.46 7.111 36.991 31.521 1.00 26.68
25	ATOM		66.010 37.176 32.464 1.00 27.90
	ATOM		65.237 35.878 32.642 1.00 25.29
	ATOM		6.511 37.697 33.810 1.00 31.23
	ATOM		5.927 38.617 34.378 1.00 37.67
20	ATOM		7.596 37.114 34.316 1.00 34.15
30	ATOM		58.174 37.550 35.588 1.00 37.23
	ATOM ATOM		59.294 36.605 36.027 1.00 40.21
	ATOM		58.785 35.324 36.361 1.00 53.99
	ATOM		38.727 38.958 35.417 1.00 33.67
35	ATOM		8.532 39.827 36.268 1.00 40.73
33	ATOM		9.411 39.171 34.298 1.00 29.95
	ATOM		70.000 40.458 33.957 1.00 29.77
	ATOM		70.684 40.350 32.594 1.00 30.79
	ATOM		71.481 41.558 32.167 1.00 31.34
40	ATOM		72.781 41.638 32.918 1.00 33.62
+0	ATOM		73.657 42.660 32.358 1.00 41.68
	ATOM		74.584 43.310 33.052 1.00 41.20
	ATOM	1750 CZ ARG 384	74.756 43.047 34.339 1.00 42.11
	ATOM	1751 NH1 ARG 384	75.349 44.213 32.455 1.00 37.27
45	ATOM		88.910 41.536 33.911 1.00 35.72
,,,	ATOM		9.090 42.635 34.439 1.00 41.66
	ATOM		7.768 41.196 33.318 1.00 34.30
	ATOM		66.646 42.119 33.199 1.00 32.40
	ATOM		55.527 41.502 32.356 1.00 29.02
50	ATOM		64.344 42.407 32.163 1.00 26.56

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ATOM 1759 CD1 PHE 385
                                64.317 43.320 31.119 1.00 26.59
    ATOM 1760 CD2 PHE 385
                                63.263 42.355 33.037 1.00 24.69
    ATOM 1761 CE1 PHE 385
                                63.231 44.173 30.947 1.00 31.70
                                62.174 43.202 32.875 1.00 26.79
    ATOM 1762 CE2 PHE 385
    ATOM 1763 CZ PHE 385
                               62.158 44.115 31.827 1.00 31.59
    ATOM 1764 C PHE 385
                               66.121 42.492 34.578 1.00 32.98
    ATOM 1765 O PHE 385
                               65.822 43.659 34.839 1.00 33.91
                               66.003 41.499 35.456 1.00 33.91
    ATOM 1766 N LEU 386
                                65.533 41.736 36.818 1.00 38.66
    ATOM 1767 CA LEU 386
    ATOM 1768 CB LEU 386
                                65.547 40.440 37.633 1.00 43.79
10
    ATOM 1769 CG LEU 386
                                64.327 39.521 37.525 1.00 49.81
    ATOM 1770 CD1 LEU 386
                                64.652 38.147 38.099 1.00 51.12
    ATOM 1771 CD2 LEU 386
                                63.135 40.148 38.246 1.00 49.17
                               66.445 42.761 37.475 1.00 38.95
    ATOM 1772 C LEU 386
    ATOM 1773 O LEU 386
                               65.979 43.682 38.146 1.00 42.16
15
    ATOM 1774 N HIS 387
                              67.745 42.613 37.248 1.00 33.62
                               68.723 43.531 37.808 1.00 39.73
    ATOM 1775 CA HIS 387
    ATOM 1776 CB HIS 387
                               70.138 42.980 37.639 1.00 40.71
    ATOM 1777 CG HIS 387
                               70.403 41.749 38.449 1.00 52.03
                                69.573 40.967 39.181 1.00 53.85
    ATOM 1778 CD2 HIS 387
20
                                71.657 41.189 38.566 1.00 54.79
    ATOM 1779 ND1 HIS 387
                                71.590 40.114 39.334 1.00 56.55
    ATOM 1780 CE1 HIS 387
    ATOM 1781 NE2 HIS 387
                                70.336 39.958 39.720 1.00 57.48
    ATOM 1782 C HIS 387
                              68.594 44.913 37.175 1.00 42.08
                              68.712 45.926 37.865 1.00 44.12
    ATOM 1783 O HIS 387
25
                               68.318 44.957 35.874 1.00 42.38
    ATOM 1784 N MET 388
                                68.154 46.229 35.175 1.00 38.00
    ATOM 1785 CA MET 388
    ATOM 1786 CB MET 388
                                67.840 46.006 33.692 1.00 40.21
    ATOM 1787 CG MET 388
                                69.009 45.555 32.829 1.00 41.26
    ATOM 1788 SD MET 388
                                68,500 45,427 31,089 1,00 45,51
30
    ATOM 1789 CE MET 388
                                69.089 43.802 30.645 1.00 42.40
    ATOM 1790 C MET 388
                               67.025 47.044 35.810 1.00 38.11
                               67.155 48.255 35.997 1.00 38.41
    ATOM 1791 O MET 388
    ATOM 1792 N LYS 389
                               65.926 46.374 36.144 1.00 39.67
                                64.773 47.036 36.750 1.00 44.96
35
    ATOM 1793 CA LYS 389
                                63.570 46.087 36.818 1.00 49.52
    ATOM 1794 CB LYS 389
    ATOM 1795 CG LYS 389
                                62.674 46.102 35.588 1.00 56.74
    ATOM 1796 CD LYS 389
                                62.145 47.509 35.278 1.00 68.05
    ATOM 1797 CE LYS 389
                                61.287 48.100 36.403 1.00 71.47
40
    ATOM 1798 NZ LYS 389
                                60.038 47.330 36.661 1.00 71.98
    ATOM 1799 C LYS 389
                               65.041 47.604 38.141 1.00 46.60
                               64.516 48.661 38.499 1.00 47.25
    ATOM 1800 O LYS 389
    ATOM 1801 N VAL 390
                               65.832 46.893 38.935 1.00 47.15
    ATOM 1802 CA VAL 390
                                66.129 47.353 40.284 1.00 50.75
    ATOM 1803 CB VAL 390
                                66,686 46,202 41,182 1,00 50,42
45
    ATOM 1804 CG1 VAL 390
                                68.095 45.802 40.770 1.00 47.93
                                66.650 46.612 42.640 1.00 56.67
    ATOM 1805 CG2 VAL 390
    ATOM 1806 C VAL 390
                               67.072 48.558 40.286 1.00 49.82
    ATOM 1807 O VAL 390
                               66.971 49.426 41.152 1.00 52.44
    ATOM 1808 N GLU 391
                               67.926 48.651 39.272 1.00 46.14
50
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	ATOM	1809 CA GLU 391	68.888 49.741 39.173 1.00 43.84
	ATOM	1810 CB GLU 391	70.150 49.268 38.449 1.00 41.44
	ATOM	1811 CG GLU 391	70.837 48.074 39.095 1.00 51.12
	ATOM	1812 CD GLU 391	71.218 48.325 40.540 1.00 57.29
5	ATOM	1813 OE1 GLU 391	71.970 49.287 40.802 1.00 58.15
_	ATOM	1814 OE2 GLU 391	70.764 47.559 41.416 1.00 62.51
	ATOM	1815 C GLU 391	68.386 51.015 38.501 1.00 45.94
	ATOM	1816 O GLU 391	68.567 52.114 39.033 1.00 51.14
	ATOM	1817 N CYA 392	67.727 50.872 37.354 1.00 45.84
10	ATOM	1818 CA CYA 392	67.255 52.029 36.598 1.00 41.60
	ATOM	1819 CB CYA 392	67.681 51.889 35.140 1.00 42.06
	ATOM	1820 SG CYA 392	69.452 52.008 34.968 1.00 44.47
	ATOM	1821 AS CYA 392	69.867 50.812 33.150 1.00 54.22
	ATOM	1822 C CYA 392	65.779 52.395 36.683 1.00 42.27
15	<b>ATOM</b>	1823 O CYA 392	64.937 51.564 37.029 1.00 43.91
	ATOM	1824 N PRO 393	65.451 53.674 36.414 1.00 42.79
	<b>ATOM</b>	1825 CD PRO 393	66.384 54.774 36.106 1.00 38.59
	<b>ATOM</b>	1826 CA PRO 393	64.067 54.159 36.459 1.00 44.20
	<b>ATOM</b>	1827 CB PRO 393	64.218 55.667 36.238 1.00 39.88
20	ATOM	1828 CG PRO 393	65.487 55.789 35.459 1.00 35.88
	ATOM	1829 C PRO 393	63.178 53.513 35.398 1.00 45.29
	ATOM	1830 O PRO 393	63.600 53.308 34.257 1.00 43.97
	ATOM	1831 N THR 394	61.935 53.238 35.782 1.00 48.20
	ATOM	1832 CA THR 394	60.959 52.607 34.901 1.00 53.71
25	ATOM	1833 CB THR 394	59.605 52.429 35.629 1.00 59.59
	ATOM	1834 OG1 THR 394	58.690 51.717 34.787 1.00 66.50
	ATOM	1835 CG2 THR 394	59.013 53.787 36.004 1.00 61.00
	ATOM	1836 C THR 394	60.752 53.358 33.581 1.00 51.35
	ATOM	1837 O THR 394	60.419 52.751 32.563 1.00 54.39
30	ATOM	1838 N GLU 395	61.008 54.664 33.595 1.00 47.65
	ATOM	1839 CA GLU 395	60.845 55.509 32.414 1.00 44.43
	ATOM	1840 CB GLU 395	60.988 56.978 32.804 1.00 43.85
	ATOM	1841 C GLU 395	61.788 55.175 31.250 1.00 42.93
2.5	ATOM	1842 O GLU 395 1843 N LEU 396	61.589 55.649 30.129 1.00 41.39 62.818 54.375 31.517 1.00 39.38
35	ATOM	1843 N LEU 396 1844 CA LEU 396	63.782 53.989 30.486 1.00 35.70
	ATOM	1844 CA LEU 396 1845 CB LEU 396	65.185 53.867 31.090 1.00 34.96
	ATOM	1846 CG LEU 396	65.854 55.141 31.609 1.00 36.47
	ATOM ATOM	1847 CD1 LEU 396	67.234 54.807 32.150 1.00 34.21
40			65.959 56.164 30.491 1.00 32.74
40	ATOM ATOM	1848 CD2 LEU 396 1849 C LEU 396	63.407 52.671 29.803 1.00 34.60
	ATOM	1850 O LEU 396	64.086 52.223 28.873 1.00 30.36
	ATOM	1851 N PHE 397	62.325 52.059 30.269 1.00 33.02
	ATOM	1852 CA PHE 397	61.868 50.792 29.725 1.00 33.39
45	ATOM	1853 CB PHE 397	61.615 49.782 30.852 1.00 34.30
7.5	ATOM	1854 CG PHE 397	62.834 49.439 31.665 1.00 32.62
	ATOM	1855 CD1 PHE 397	63.296 50.301 32.654 1.00 32.35
	ATOM	1856 CD2 PHE 397	63.504 48.241 31.461 1.00 31.28
	ATOM	1857 CEI PHE 397	64.407 49.976 33.426 1.00 27.01
50	ATOM	1858 CE2 PHE 397	64.616 47.905 32.229 1.00 33.34

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ATOM 1859 CZ PHE 397
                               65.067 48.775 33.213 1.00 31.29
                              60.580 50.961 28.934 1.00 33.17
    ATOM 1860 C PHE 397
                               59.540 51.318 29.498 1.00 31.99
    ATOM 1861 O PHE 397
    ATOM 1862 N PRO 398
                               60.636 50.752 27.606 1.00 32.45
    ATOM 1863 CD PRO 398
                                61.821 50.493 26.768 1.00 28.15
    ATOM 1864 CA PRO 398
                                59.429 50.885 26.786 1.00 30.02
    ATOM 1865 CB PRO 398
                               59.921 50.483 25.394 1.00 28.15
                               61.352 50.923 25.397 1.00 24.89
    ATOM 1866 CG PRO 398
                               58.384 49.900 27.326 1.00 28.39
    ATOM 1867 C PRO 398
                               58.735 48.810 27.789 1.00 28.00
    ATOM 1868 O PRO 398
10
    ATOM 1869 N PRO 399
                               57.092 50.262 27.267 1.00 32.45
                               56.577 51.511 26.672 1.00 34.93
    ATOM 1870 CD PRO 399
    ATOM 1871 CA PRO 399
                               55.989 49.421 27.753 1.00 32.54
                                54.755 50.122 27.188 1.00 34.47
    ATOM 1872 CB PRO 399
    ATOM 1873 CG PRO 399
                                55.159 51.564 27.196 1.00 31.37
15
    ATOM 1874 C PRO 399
                               56.044 47.946 27.338 1.00 32.18
                               55.950 47.054 28.188 1.00 32.58
    ATOM 1875 O PRO 399
    ATOM 1876 N LEU 400
                               56.195 47.689 26.041 1.00 30.15
                                56.259 46.314 25.541 1.00 32.32
    ATOM 1877 CA LEU 400
                                56.211 46.297 24.011 1.00 28.67
    ATOM 1878 CB LEU 400
20
                                56.028 44.927 23.351 1.00 28.77
    ATOM 1879 CG LEU 400
                               54.802 44.234 23.919 1.00 22.73
    ATOM 1880 CD1 LEU 400
    ATOM 1881 CD2 LEU 400
                               55.897 45.096 21.846 1.00 27.89
                               57.496 45.561 26.051 1.00 32.27
    ATOM 1882 C LEU 400
    ATOM 1883 O LEU 400
                               57.437 44.358 26.307 1.00 32.87
25
                               58.602 46.279 26.215 1.00 32.27
    ATOM 1884 N PHE 401
                              59.847 45.695 26.710 1.00 32.39
    ATOM 1885 CA PHE 401
                                60.946 46.769 26.711 1.00 31.38
    ATOM 1886 CB PHE 401
                                62.290 46.286 27.194 1.00 35.12
     ATOM 1887 CG PHE 401
     ATOM 1888 CD1 PHE 401
                               62.835 45.089 26.729 1.00 34.68
30
                                63.030 47.051 28.097 1.00 34.57
     ATOM 1889 CD2 PHE 401
                                64.100 44.662 27.155 1.00 30.27
    ATOM 1890 CE1 PHE 401
     ATOM 1891 CE2 PHE 401
                               64.291 46.635 28.526 1.00 33.57
                               64.828 45.438 28.054 1.00 35.74
     ATOM 1892 CZ PHE 401
    ATOM 1893 C PHE 401
                               59.599 45.169 28.129 1.00 32.21
35
                               60.002 44.056 28.478 1.00 33.36
     ATOM 1894 O PHE 401
                               58.902 45.967 28.929 1.00 31.85
     ATOM 1895 N LEU 402
                               58.582 45.602 30.302 1.00 35.06
     ATOM 1896 CA LEU 402
     ATOM 1897 CB LEU 402
                                57.948 46.789 31.029 1.00 34.76
     ATOM 1898 CG LEU 402
                                58.878 47.852 31.591 1.00 33.48
40
                                58.060 49.010 32.152 1.00 32.58
     ATOM 1899 CD1 LEU 402
     ATOM 1900 CD2 LEU 402
                                59,753 47,217 32,662 1.00 26,27
                               57.626 44.426 30.393 1.00 36.80
     ATOM 1901 C LEU 402
                               57.793 43.545 31.239 1.00 35.43
     ATOM 1902 O LEU 402
                               56.600 44.443 29.547 1.00 38.50
     ATOM 1903 N GLU 403
45
                                55.581 43.401 29.540 1.00 40.24
     ATOM 1904 CA GLU 403
                                54.435 43.792 28.605 1.00 44.03
     ATOM 1905 CB GLU 403
                                53.239 42.850 28.666 1.00 55.53
     ATOM 1906 CG GLU 403
     ATOM 1907 CD GLU 403
                                52.180 43.159 27.618 1.00 66.67
     ATOM 1908 OE1 GLU 403
                               52.151 44.299 27.095 1.00 70.81
50
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	ATOM	1909 OE2 GLU 403	51.370 42.255 27.315 1.00 73.80
	ATOM	1910 C GLU 403	56.096 42.018 29.162 1.00 38.00
	ATOM	1911 O GLU 403	55.745 41.029 29.805 1.00 38.78
	ATOM	1912 N VAL 404	56.934 41.955 28.132 1.00 37.39
5	ATOM	1913 CA VAL 404	57.475 40.686 27.652 1.00 37.05
-	ATOM	1914 CB VAL 404	58.180 40.855 26.286 1.00 35.57
	ATOM	1915 CG1 VAL 404	58.677 39.513 25.776 1.00 36.85
	ATOM	1916 CG2 VAL 404	57.222 41.451 25.287 1.00 42.03
	ATOM	1917 C VAL 404	58.438 40.000 28.609 1.00 38.69
10	ATOM	1918 O VAL 404	58.436 38.774 28.727 1.00 40.71
	<b>ATOM</b>	1919 N PHE 405	59.267 40.785 29.286 1.00 39.34
	ATOM	1920 CA PHE 405	60.250 40.221 30.198 1.00 39.33
	ATOM	1921 CB PHE 405	61.620 40.840 29.913 1.00 33.87
	ATOM	1922 CG PHE 405	62.107 40.609 28.509 1.00 32.17
15	ATOM	1923 CD1 PHE 405	62.355 41.683 27.660 1.00 31.34
	ATOM	1924 CD2 PHE 405	62.315 39.317 28.032 1.00 31.98
	ATOM	1925 CE1 PHE 405	62.801 41.476 26.352 1.00 30.79
	ATOM	1926 CE2 PHE 405	62.759 39.099 26.730 1.00 26.06
	<b>ATOM</b>	1927 CZ PHE 405	63.004 40.182 25.889 1.00 27.98
20	ATOM	1928 C PHE 405	59.905 40.322 31.682 1.00 42.64
	ATOM	1929 O PHE 405	60.785 40.188 32.534 1.00 45.10
	ATOM	1930 N GLU 406	58.630 40.536 31.988 1.00 48.95
	ATOM	1931 CA GLU 406	58.181 40.641 33.373 1.00 56.93
	ATOM	1932 CB GLU 406	56.820 41.324 33.432 1.00 56.94
25	ATOM	1933 C GLU 406	58.116 39.263 34.040 1.00 61.92
	ATOM	1934 O GLU 406	57.988 38.256 33.308 1.00 67.61
	ATOM	1 O1 HOH 501	67.588 36.828 11.225 1.00 27.32
	ATOM	2 O1 HOH 502	68.647 41.203 12.940 1.00 39.54
	ATOM	3 O1 HOH 503	64.072 40.115 12.407 1.00 32.47
30	ATOM	4 O1 HOH 504	62.312 39.659 16.075 1.00 17.39
	ATOM	5 O1 HOH 505	63,449 46,468 15,530 1.00 30,46
	ATOM	6 O1 HOH 506	67.191 15.561 -0.279 1.00 35.96
	ATOM	7 O1 HOH 507	67.100 11.855 0.295 1.00 20.00
	ATOM	8 O1 HOH 508	61.004 15.510 0.047 1.00 20.00
35	ATOM	9 O1 HOH 509	59.851 10.761 6.050 1.00 20.00
	ATOM	10 O1 HOH 510	57.553 11.824 10.360 1.00 44.63 54.101 13.545 8.720 1.00 20.00
	ATOM	11 O1 HOH 511	• ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
	ATOM	12 O1 HOH 512	=
40	ATOM	13 O1 HOH 513	
40	ATOM	14 O1 HOH 514	50.474 22.912 7.942 1.00 45.34 49.737 20.631 11.530 1.00 20.00
	ATOM	15 O1 HOH 515	
	ATOM	16 O1 HOH 516	
	ATOM	17 O1 HOH 517	53.818 25.833 10.682 1.00 42.12 52.591 31.216 7.313 1.00 35.55
4.5	ATOM	18 O1 HOH 518	58.510 31.667 2.158 1.00 20.00
45	ATOM	19 O1 HOH 519	58.235 36.751 2.232 1.00 20.00
	ATOM	20 O1 HOH 520	62.484 37.992 5.537 1.00 20.00
	ATOM		68.184 36.969 5.889 1.00 50.08
	ATOM		66.889 33.781 8.584 1.00 20.00
50	ATOM		67.217 30.836 3.085 1.00 34.44
50	ATOM	24 OI HON 324	07.217 30.030 3.003 1.00 5 1.11

	ATOM	25 O1 HOH	525	64.336 28.325 3.098 1.00 20.00
	ATOM	26 O1 HOH	526	67.667 26.625 1.519 1.00 20.00
	ATOM	27 O1 HOH	527	76.757 22.883 5.467 1.00 36.94
	ATOM	28 O1 HOH	528	72.250 17.936 6.950 1.00 36.00
5	ATOM	29 O1 HOH		71.760 14.791 8.058 1.00 40.18
•	ATOM	30 O1 HOH		72.884 14.751 11.484 1.00 41.44
	ATOM	31 O1 HOH		69.235 12.986 11.709 1.00 39.38
	ATOM	32 O1 HOH		69.402 12.036 14.891 1.00 40.68
	ATOM	33 O1 HOH		64.560 10.910 15.076 1.00 20.00
10	ATOM	34 O1 HOH		63.169 10.413 11.722 1.00 20.00
10	ATOM	35 O1 HOH		66.042 11.455 11.077 1.00 41.05
	ATOM	36 O1 HOH		76.285 12.458 10.677 1.00 20.00
	ATOM	37 O1 HOH		81.094 22.520 13.435 1.00 48.70
	ATOM	38 O1 HOH		80.505 25.457 14.849 1.00 46.30
15	ATOM	39 O1 HOH		77.669 21.932 18.119 1.00 43.79
13	ATOM	40 O1 HOH		77.187 28.903 21.137 1.00 40.22
	ATOM	41 O1 HOH		76.420 30.760 23.658 1.00 29.63
	ATOM	42 O1 HOH		83.028 32.743 20.922 1.00 38.14
		43 O1 HOH		82.842 43.133 17.983 1.00 39.36
20	ATOM	44 O1 HOH		77.484 34.040 9.664 1.00 36.37
20	ATOM	45 O1 HOH		75.904 32.986 12.256 1.00 34.93
	ATOM			74.185 29.689 9.761 1.00 38.60
	ATOM			64.936 20.644 23.365 1.00 36.83
	ATOM	• • • • • • • • • • • • • • • • • • • •		61.750 22.313 25.288 1.00 34.81
25	ATOM			59.544 21.463 26.162 1.00 20.00
25	ATOM			62,300 27.528 24.386 1.00 35.89
	ATOM	50 O1 HOH 51 O1 HOH		58.228 29.424 24.603 1.00 25.47
	ATOM	52 O1 HOH		57.368 32.196 30.527 1.00 45.27
	ATOM ATOM	53 O1 HOH		62.063 36.304 30.245 1.00 42.26
20		54 O1 HOH		64.722 36.725 28.906 1.00 24.66
30	ATOM			62.207 35.851 26.642 1.00 30.36
	ATOM			63.608 33.715 25.707 1.00 42.74
	ATOM	56 O1 HOH 57 O1 HOH		62.979 38.422 32.977 1.00 42.74
	ATOM			66.911 33.364 34.901 1.00 50.02
25	ATOM			72.608 29.636 31.674 1.00 37.60
35	ATOM	59 O1 HOH		76.967 40.633 32.514 1.00 44.81
	ATOM	60 O1 HOH 61 O1 HOH		73.613 41.817 36.847 1.00 31.79
	ATOM			75.773 46.227 30.514 1.00 29.06
	ATOM			79.903 46.178 30.800 1.00 41.67
40	ATOM			69.746 51.175 33.564 1.00 20.00
40	ATOM	64 O1 HOH		74.320 52.047 39.438 1.00 20.00
	ATOM	65 O1 HOH		65.900 53.647 27.404 1.00 40.45
	ATOM	66 O1 HOH		
	ATOM	67 O1 HOH		68.848 53.076 17.895 1.00 39.25 63.507 48.672 13.581 1.00 43.77
4.5	ATOM	68 O1 HOH		
45	ATOM	69 O1 HOH		64.625 46.825 10.331 1.00 20.00
	ATOM	70 O1 HOH		55.882 41.431 11.148 1.00 20.00 52.830 43.513 20.032 1.00 35.18
	ATOM	71 O1 HOH		
	ATOM	72 O1 HOH		56.990 49.485 24.052 1.00 37.30 54.188 47.024 30.900 1.00 52.93
<b>50</b>	ATOM	73 O1 HOH		
50	ATOM	74 O1 HOH	574	57.823 44.590 34.025 1.00 53.64

	ATOM	75 O1 HOH 575	47.827 29.597 30.690 1.00 37.61
	ATOM		53.030 24.901 32.732 1.00 45.06
	ATOM	76 O1 HOH 576	
	ATOM	77 O1 HOH 577	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
_	ATOM	78 O1 HOH 578	17.232 201242 20144
5	ATOM	79 O1 HOH 579	<b>51.</b> 700 <b>5</b> 51
	ATOM	80 O1 HOH 580	52.831 23.395 1.634 1.00 20.00
	ATOM	81 O1 HOH 581	51.472 22.968 -0.900 1.00 25.10
	ATOM	82 O1 HOH 582	77.238 52,503 8.906 1.00 47.05
	END		
10	ATOM	2004 C1 DMT 1	67.320 42.326 18.648 1.00 28.58
	ATOM	2005 C2 DMT 1	68.927 43.263 23.318 1.00 29.26
	ATOM	2006 C3 DMT 1	67.236 43.583 19.236 1.00 24.54
	ATOM	2007 C4 DMT 1	69.268 44.313 24.111 1.00 28.48
	ATOM	2008 C5 DMT 1	68.003 43.859 20.363 1.00 28.76
15	<b>ATOM</b>	2009 C6 DMT 1	68.654 44.389 25.458 1.00 28.16
	ATOM	2010 C7 DMT 1	68.811 42.902 20.875 1.00 26.80
	ATOM	2011 C8 DMT 1	67.803 43.410 25.793 1.00 29.83
	ATOM	2012 C9 DMT 1	68.921 41.665 20.324 1.00 26.77
	<b>ATOM</b>	2013 C10 DMT 1	67.464 42.358 24.989 1.00 28.60
20	ATOM	2014 C11 DMT 1	68.165 41.349 19.185 1.00 25.29
	ATOM	2015 C12 DMT 1	68.059 42.281 23.675 1.00 26.74
	<b>ATOM</b>	2016 C13 DMT 1	66.475 42.038 17.456 1.00 21.51
	ATOM	2017 C14 DMT 1	68.916 45.478 26.380 1.00 21.05
	ATOM	2018 C15 DMT 1	66.989 40.910 16.417 1.00 22.84
25	ATOM	2019 C16 DMT 1	68.090 46.870 26.009 1.00 19.41
	ATOM	2020 C17 DMT 1	65.982 40.730 15.243 1.00 27.07
	ATOM	2021 C18 DMT 1	70.279 46.131 26.085 1.00 16.03
	ATOM	2022 C19 DMT 1	67.903 45.249 20.974 1.00 19.56
	ATOM	2023 C20 DMT 1	69.853 40.599 20.901 1.00 4.52
30	ATOM	2024 N1 DMT 1	68.280 41.070 16.042 1.00 17.57
	ATOM	2025 O1 DMT 1	67.209 43.465 27.087 1.00 25.94
	ATOM	2026 O2 DMT 1	69.547 43.191 22.015 1.00 30.23
	ATOM	2027 O3 DMT 1	66.449 40.778 14.118 1.00 29.45
	ATOM	2028 O4 DMT 1	64.820 40.564 15.546 1.00 26.46
35	END		

WO 99/26966

PCT/US98/25296

#### APPENDIX 4

#### TR TRIAC.PDB

**REMARK** 

REMARK TR triac full length numbering

REMARK Rfactor 0.236 Rfree 0.241

REMARK Resolution 25. 2.5 all reflections

REMARK

REMARK Three cacodylate-modified cysteines:

REMARK Cys334, Cys380, Cys392

REMARK modeled as free arsenic atoms 10

REMARK

REMARK conserved polar HOH numbered as in TR\_t3.pdb

REMARK rearrangements start 600

REMARK side chain of certain residues modeled as ALA due to poor density; 15

REMARK however, residue name reflects true residue for clarity

REMARK

REMARK clone obtained from Murray et. al.

REMARK deposited sequence confirmed,

REMARK differing from that reported by Thompson et. al. 20

REMARK in the following codons:

REMARK 281 Thr - Ala

REMARK 285 Lys - Glu

REMARK identical to that reported by Mitsuhashi et. al.

REMARK gb:RNTRAVI X07409 25

M.B. MURRAY, N.D.ZILZ, AUTH **JRNL** 

N.L.MCCREARY, M.J.MACDONALD

JRNL **AUTH 2 H.C.TOWLE** 

TITL ISOLATION AND CHARACTERIZATION OF RAT CDNA JRNL

**CLONES FOR TWO** 30

JRNL

45

TITL 2 DISTINCT THYROID HORMONE RECPTORS JRNL

**JRNL** REF JBC

V. 263 25 1988 AUTH C.C.THOMPSON, C.WEINBERGER, R.LEBO, R.M.EVANS

V. 237 1987

IDENTIFICATION OF A NOVEL THYROID HORMONE JRNL

RECEPTOR EXPRESSED 35

TITL 2 IN THE MAMMALIAN CENTRAL NERVOUS SYSTEM JRNL

**JRNL** REF SCIENCE

AUTH T.MITSUHASHI,G.TENNYSON,V.NIKODEM **JRNL** 

TITL NUCLEOTIDE SEQUENCE OF NOVEL CDNAS GENERATED JRNL

40 BY ALTERNATIVE

TITL 2 SPLICING OF A RAT THYROID HORMONE RECEPTOR **JRNL** 

GENE TRANSCRIPT REF NUC. ACIDS. RES. V. 16 12 1988 JRNL

REMARK 9.880 -24.199 7.196 1.00 57.79 ATOM 1 CB ARG 157

11.380 -24.411 7.340 1.00 57.79 ATOM 2 CG ARG 157

11.960 -23.602 8.486 1.00 57.79 3 CD ARG 157 ATOM

11.492 -24.098 9.778 1.00 57.79 ATOM 4 NE ARG 157

	ATOM	5 CZ ARG 157	12.284 -24.379 10.809 1.00 57.79
	ATOM	6 NH1 ARG 157	13.598 -24.212 10.714 1.00 57.79
	ATOM	7 NH2 ARG 157	11.762 -24.854 11.932 1.00 57.79
	ATOM	8 C ARG 157	7.774 -24.838 5.974 1.00 38.50
5	ATOM	9 O ARG 157	7.553 -24.416 4.840 1.00 57.79
-	ATOM	10 N ARG 157	9,929 -25,500 5.089 1.00 38.50
	ATOM	11 CA ARG 157	9.183 -25.276 6.360 1.00 38.50
	ATOM	12 N PRO 158	6.802 -24.951 6.895 1.00 23.08
	ATOM	13 CD PRO 158	6.945 -25.424 8.282 1.00 28.38
10	ATOM	14 CA PRO 158	5.415 -24.562 6.617 1.00 23.08
10	ATOM	15 CB PRO 158	4.704 -24.824 7.948 1.00 28.38
	ATOM	16 CG PRO 158	5.801 -24.735 8.966 1.00 28.38
	ATOM	17 C PRO 158	5.210 -23.124 6.132 1.00 23.08
	ATOM	18 O PRO 158	5.678 -22.167 6.753 1.00 28.38
16		19 N GLU 159	4.504 -23.000 5.012 1.00 19.26
15	ATOM		4.191 -21.717 4.389 1.00 19.26
	ATOM		4.022 -21.912 2.878 1.00 24.58
	ATOM		5.317 -22.009 2.086 1.00 24.58
	ATOM		5.849 -20.651 1.659 1.00 24.58
	ATOM	23 CD GLU 159	5.034 -19.722 1.476 1.00 24.58
20	ATOM	24 OE1 GLU 159	7.080 -20.513 1.490 1.00 24.58
	ATOM	25 OE2 GLU 159	2.879 -21.193 4.968 1.00 19.26
	ATOM	26 C GLU 159	2.152 -21.931 5.636 1.00 24.58
	ATOM	27 O GLU 159	2.579 -19.899 4.765 1.00 17.44
	ATOM	28 N PRO 160	
25	ATOM	29 CD PRO 160	
	ATOM	30 CA PRO 160	
	ATOM	31 CB PRO 160	
	ATOM	32 CG PRO 160	
	ATOM	33 C PRO 160	0.098 -20.006
30	ATOM	34 O PRO 160	
	ATOM	35 N THR 161	* · - · -
	ATOM	36 CA THR 161	
	ATOM	37 CB THR 161	
	ATOM	38 OG1 THR 161	<del>- • • • • • • • • • • • • • • • • • • •</del>
35	ATOM	39 CG2 THR 161	· · · · · · · · · · · · · · · · · · ·
	ATOM	40 C THR 161	
	ATOM	41 O THR 161	2.071
	ATOM	42 N PRO 162	-3.918 -20.200 3.449 1.00 12.94
	ATOM	43 CD PRO 162	-4.311 -21.559 3.038 1.00 17.56
40	ATOM	44 CA PRO 162	-4.743 -19.190 2.780 1.00 12.94
	ATOM	45 CB PRO 162	-5.846 -20.029 2.143 1.00 17.56
	ATOM	46 CG PRO 162	-5.147 -21.303 1.816 1.00 17.56
	ATOM	47 C PRO 162	-5.317 -18.171 3.763 1.00 12.94
	ATOM	48 O PRO 162	-5.305 -16.964 3.503 1.00 17.56
45	ATOM	49 N GLU 163	-5.790 -18.668 4.903 1.00 19.45
	ATOM	50 CA GLU 163	-6.374 -17.828 5.943 1.00 19.45
	ATOM	51 CB GLU 163	-6.994 -18.690 7.047 1.00 49.96
	ATOM	52 CG GLU 163	-8.178 -19.558 6.606 1.00 49.96
	ATOM	53 CD GLU 163	-7.782 -20.720 5.697 1.00 49.96
50	ATOM	54 OE1 GLU 163	-6.735 -21.361 5.951 1.00 49.96

WO 99/26966

WO 99/26966

### PCT/US98/25296

	ATOM	55 OE2 GLU 163	-8,527 -20.999 4.731 1.00 49.96
	ATOM	56 C GLU 163	-5.330 -16.897 6.548 1.00 19.45
	ATOM	57 O GLU 163	-5.614 -15.731 6.832 1.00 49.96
	ATOM	58 N GLU 164	-4.120 -17.417 6.734 1.00 22.03
5	ATOM	59 CA GLU 164	-3.033 -16.634 7.305 1.00 22.03
,	ATOM	60 CB GLU 164	-1.875 -17.541 7.725 1.00 17.15
	ATOM	61 CG GLU 164	-2.198 -18.414 8.937 1.00 17.15
	ATOM	62 CD GLU 164	-1.114 -19.434 9.249 1.00 17.15
	ATOM	63 OE1 GLU 164	-0.283 -19.710 8.361 1.00 17.15
10	ATOM	64 OE2 GLU 164	-1.099 -19.968 10.379 1.00 17.15
10	ATOM	65 C GLU 164	-2.559 -15.542 6.354 1.00 22.03
	ATOM	66 O GLU 164	-2.160 -14.470 6.802 1.00 17.15
	ATOM	67 N TRP 165	-2.607 -15.805 5.048 1.00 10.72
	ATOM	68 CA TRP 165	-2.205 -14.803 4.063 1.00 10.72
15	ATOM	69 CB TRP 165	-2.223 -15.377 2.644 1.00 2.00
15	ATOM	70 CG TRP 165	-0.928 -16.003 2.227 1.00 2.00
	ATOM	71 CD2 TRP 165	0.350 -15.358 2.131 1.00 2.00
	ATOM	72 CE2 TRP 165	1.275 -16.326 1.685 1.00 2.00
		73 CE3 TRP 165	0.804 -14.054 2.379 1.00 2.00
20	ATOM	74 CD1 TRP 165	-0.731 -17.298 1.848 1.00 2.00
20	ATOM ATOM	75 NE1 TRP 165	0.587 -17.500 1.521 1.00 2.00
	ATOM	76 CZ2 TRP 165	2.627 -16.036 1.479 1.00 2.00
	ATOM	77 CZ3 TRP 165	2.152 -13.764 2.174 1.00 2.00
	ATOM	78 CH2 TRP 165	3.046 -14.754 1.729 1.00 2.00
25	ATOM	79 C TRP 165	-3.137 -13.601 4.149 1.00 10.72
23	ATOM	80 O TRP 165	-2.717 -12.463 3.925 1.00 2.00
	ATOM	81 N ASP 166	-4.408 -13.861 4.441 1.00 14.80
	ATOM	82 CA ASP 166	-5.397 -12.796 4.580 1.00 14.80
	ATOM	83 CB ASP 166	-6.812 -13.370 4.698 1.00 28.74
30	ATOM	84 CG ASP 166	-7.298 -13.999 3.403 1.00 28.74
50	ATOM	85 OD1 ASP 166	-6.909 -13.511 2.320 1.00 28.74
	ATOM	86 OD2 ASP 166	-8.071 -14.978 3.466 1.00 28.74
	ATOM	87 C ASP 166	-5.063 -11.981 5.819 1.00 14.80
	ATOM	88 O ASP 166	-5.056 -10.749 5.775 1.00 28.74
35	ATOM	89 N LEU 167	-4.745 -12.682 6.906 1.00 11.01
55	ATOM	90 CA LEU 167	-4.383 -12.044 8.166 1.00 11.01
	ATOM	91 CB LEU 167	-4.036 -13.103 9.214 1.00 31.53
	ATOM	92 CG LEU 167	-4.672 -12.975 10.601 1.00 31.53
	ATOM	93 CD1 LEU 167	-3.806 -13.709 11.619 1.00 31.53
40	ATOM	94 CD2 LEU 167	-4.820 -11.507 10.989 1.00 31.53
70	ATOM	95 C LEU 167	-3.161 -11.159 7.933 1.00 11.01
	ATOM	96 O LEU 167	-3.120 -10.006 8.367 1.00 31.53
	ATOM	97 N ILE 168	-2.180 -11.714 7.228 1.00 13.18
	ATOM	98 CA ILE 168	-0.937 -11.027 6.900 1.00 13.18
45	ATOM	99 CB ILE 168	0.015 -11.968 6.113 1.00 18.30
73	ATOM	100 CG2 ILE 168	1.118 -11.182 5.414 1.00 18.30
	ATOM	100 CG2 ILE 108	0.604 -13.013 7.063 1.00 18.30
	ATOM	101 COT ILE 108	1.379 -14.111 6.373 1.00 18.30
	ATOM	102 CD1 ILE 108	-1.185 -9.747 6.107 1.00 13.18
50	ATOM	103 C ILE 108	-0.637 -8.697 6.437 1.00 18.30
50	AIOM	104 O ILL 100	0.057 -0.077 0.157 1.00 20.50

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-2.032 -9.831 5.084 1.00 12.99
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            106 CA HIS 169
     ATOM
                               -3.218 -9.087 3.062 1.00 13.09
            107 CB HIS 169
     ATOM
                               -2.553 -10.045 2.126 1.00 13.09
            108 CG HIS 169
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            109 CD2 HIS 169
     ATOM
                                -3.249 -11.000 1.416 1.00 13.09
            110 ND1 HIS 169
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                               -2.403 -11.728 0.710 1.00 13.09
            111 CE1 HIS 169
    ATOM
            112 NE2 HIS 169
                               -1.181 -11.277 0.936 1.00 13.09
     ATOM
                              -3.017 -7.550 5.017 1.00 12.99
     ATOM
            113 C HIS 169
            114 O HIS 169
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10
    ATOM
            115 N VAL 170
                               -3.978 -7.909 5.862 1.00 13.36
     ATOM
                                -4.696 -6.926 6.664 1.00 13.36
     ATOM
            116 CA VAL 170
     ATOM
            117 CB VAL 170
                                -5.863 -7.572 7.443 1.00 20.12
             118 CG1 VAL 170
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     ATOM
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            119 CG2 VAL 170
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     ATOM
            121 O VAL 170
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     ATOM
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                               -2.920 -7.043 8.320 1.00 11.04
     ATOM
     ATOM
            123 CA ALA 171
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            124 CB ALA 171
                                -1.249 -7.653 10.005 1.00 13.43
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            125 C ALA 171
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     ATOM
                               -0.658 -4.507 9.058 1.00 13.43
            126 O ALA 171
     ATOM
            127 N THR 172
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     ATOM
                                 0.606 -5.301 6.723 1.00 12.51
            128 CA THR 172
     ATOM
                                 1.062 -6.032 5.445 1.00 14.17
            129 CB THR 172
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     ATOM
     ATOM
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            130 OG1 THR 172
                                 2.175 -5.255 4.756 1.00 14.17
            131 CG2 THR 172
     ATOM
                                0.045 -3.936 6.337 1.00 12.51
             132 C THR 172
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                                0.701 -2.910 6.537 1.00 14.17
            133 O THR 172
     ATOM
                                -1.178 -3.921 5.815 1.00 17.79
             134 N GLU 173
     ATOM
30
                                -1.818 -2.675 5.421 1.00 17.79
             135 CA GLU 173
     ATOM
                                -3.130 -2.946 4.682 1.00 49.44
             136 CB GLU 173
     ATOM
                                -3.823 -1.679 4.171 1.00 49.44
             137 CG GLU 173
     ATOM
                                 -2.930 -0.835 3.266 1.00 49.44
            138 CD GLU 173
     ATOM
            139 OE1 GLU 173
                                 -2.075 -1.408 2.552 1.00 49.44
35
     ATOM
                                 -3.085 0.404 3.269 1.00 49.44
            140 OE2 GLU 173
     ATOM
            141 C GLU 173
                               -2.072 -1.780 6.628 1.00 17.79
     ATOM
                                -1.854 -0.568 6.557 1.00 49.44
             142 O GLU 173
     ATOM
                                -2.525 -2.375 7.731 1.00 13.12
             143 N ALA 174
     ATOM
                                -2.798 -1.631 8.957 1.00 13.12
             144 CA ALA 174
40
     ATOM
                                -3.226 -2.576 10.068 1.00 17.51
             145 CB ALA 174
     ATOM
                                -1.556 -0.856 9.375 1.00 13.12
             146 C ALA 174
     ATOM
                                -1.634 0.319 9.735 1.00 17.51
     ATOM
             147 O ALA 174
             148 N HIS 175
                               -0.409 -1.521 9.317 1.00 12.20
     ATOM
             149 CA HIS 175
                                0.851 -0.895 9.679 1.00 12.20
45
     ATOM
                                1.944 -1.949 9.886 1.00 17.52
             150 CB HIS 175
     ATOM
                                3.302 -1.365 10.136 1.00 17.52
             151 CG HIS 175
     ATOM
                                 3.733 -0.468 11.055 1.00 17.52
     ATOM
             152 CD2 HIS 175
                                 4.400 -1.679 9.364 1.00 17.52
     ATOM
             153 ND1 HIS 175
                                 5.447 -0.999 9.793 1.00 17.52
     ATOM
             154 CEI HIS 175
50
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		155 NEO HIG. 175	5.070 -0.258 10.818 1.00 17.52
	ATOM	155 NE2 HIS 175	
	ATOM	156 C HIS 175	1.311 0.133 8.654 1.00 12.20
	ATOM	157 O HIS 175	1.700 1.240 9.024 1.00 17.52
	ATOM	158 N ARG 176	1.291 -0.233 7.375 1.00 12.54
5	ATOM	159 CA ARG 176	1.735 0.677 6.328 1.00 12.54
	ATOM	160 CB ARG 176	1.662 0.017 4.950 1.00 50.41
	ATOM	161 CG ARG 176	2.683 -1.088 4.730 1.00 50.41
	ATOM	162 CD ARG 176	2.666 -1.565 3.299 1.00 50.41
	ATOM	163 NE ARG 176	3.682 -2.571 2.989 1.00 50.41
10	ATOM	164 CZ ARG 176	3.577 -3.472 2.012 1.00 50.41
	ATOM	165 NH1 ARG 176	2.496 -3.513 1.236 1.00 50.41
	<b>ATOM</b>	166 NH2 ARG 176	4.536 -4.376 1.841 1.00 50.41
	<b>ATOM</b>	167 C ARG 176	0.972 1.988 6.306 1.00 12.54
	ATOM	168 O ARG 176	1.561 3.040 6.087 1.00 50.41
15	ATOM	169 N SER 177	-0.326 1.935 6.581 1.00 24.74
	ATOM	170 CA SER 177	-1.147 3.145 6.584 1.00 24.74
	ATOM	171 CB SER 177	-2.622 2.792 6.414 1.00 21.56
	ATOM	172 OG SER 177	-3.069 1.913 7.436 1.00 21.56
	ATOM	173 C SER 177	-0.960 4.013 7.832 1.00 24.74
20	ATOM	174 O SER 177	-1.401 5.159 7.863 1.00 21.56
20	ATOM	174 O BER 177	-0.347 3.453 8.870 1.00 17.96
	ATOM	176 CA THR 178	-0.104 4.181 10.115 1.00 17.96
	ATOM	170 CA THR 178	-0.736 3.440 11.323 1.00 19.76
	ATOM	177 CB THR 178	-0.265 2.091 11.361 1.00 19.76
25	ATOM	178 OG1 THR 178	-2.253 3.443 11.211 1.00 19.76
23	ATOM	180 C THR 178	1.376 4.395 10.382 1.00 17.96
	ATOM	181 O THR 178	1.760 4.880 11.445 1.00 19.76
	ATOM	182 N ASN 179	2.207 4.024 9.417 1.00 25.88
		183 CA ASN 179	3.654 4.180 9.546 1.00 25.88
20	ATOM		4.362 2.974 8.943 1.00 44.29
30	ATOM	184 CB ASN 179	
	ATOM	185 CG ASN 179	·
	ATOM	186 OD1 ASN 179	
	ATOM	187 ND2 ASN 179	
	ATOM	188 C ASN 179	
35	ATOM	189 O ASN 179	
	ATOM	190 N ALA 180	4.332 6.502 9.604 1.00 45.20
	ATOM	191 CA ALA 180	4.740 7.818 9.126 1.00 45.20
	ATOM	192 CB ALA 180	5.026 8.743 10.313 1.00 36.14
	ATOM	193 C ALA 180	5.931 7.808 8.170 1.00 45.20
40	ATOM	194 O ALA 180	6.918 7.097 8.372 1.00 36.14
	ATOM	195 N ALA 181	5.784 8.552 7.080 1.00 44.05
	ATOM	196 CA ALA 181	6.834 8.661 6.072 1.00 44.05
	ATOM	197 CB ALA 181	8.170 9.116 6.722 1.00 50.21
	ATOM	198 C ALA 181	7.069 7.427 5.196 1.00 44.05
45	ATOM	199 O ALA 181	7.663 7.550 4.118 1.00 50.21
	ATOM	200 N GLY 182	6.567 6.268 5.622 1.00 39.06
	<b>ATOM</b>	201 CA GLY 182	6.756 5.040 4.867 1.00 39.06
	ATOM	202 C GLY 182	8.202 4.769 4.482 1.00 39.06
	ATOM	203 O GLY 182	9.096 4.785 5.334 1.00 48.58
50	ATOM	204 N SER 183	8.438 4.564 3.189 1.00 64.55

	ATOM	205 CA SER 183	9.781 4.270 2.693 1.00 64.55
	ATOM	206 CB SER 183	9.690 3.402 1.430 1.00 67.68
	<b>ATOM</b>	207 OG SER 183	8.822 3.978 0.467 1.00 67.68
	ATOM	208 C SER 183	10.643 5.510 2.437 1.00 64.55
5	ATOM	209 O SER 183	11.839 5.407 2.158 1.00 67.68
	ATOM	210 N HIS 184	10.035 6.683 2.579 1.00 52.73
	ATOM	211 CA HIS 184	10.725 7.953 2.352 1.00 52.73
	ATOM	212 CB HIS 184	9.772 8.955 1.698 1.00 44.77
	ATOM	213 C HIS 184	11.364 8.582 3.595 1.00 52.73
10	ATOM	214 O HIS 184	11.837 9.722 3.540 1.00 44.77
	ATOM	215 N TRP 185	11.420 7.842 4.699 1.00 54.14
	ATOM	216 CA TRP 185	11.977 8.389 5.940 1.00 54.14
	ATOM	217 CB TRP 185	11.813 7.395 7.104 1.00 40.24
	ATOM	218 CG TRP 185	12.605 6.123 6.991 1.00 40.24
15	ATOM	219 CD2 TRP 185	13.894 5.873 7.551 1.00 40.24
13	ATOM	220 CE2 TRP 185	14.245 4.543 7.221 1.00 40.24
		221 CE3 TRP 185	14.791 6.641 8.300 1.00 40.24
	ATOM		12.227 4.973 6.359 1.00 40.24
	ATOM		13.210 4.015 6.496 1.00 40.24
20	ATOM		15.461 3.968 7.619 1.00 40.24
20	ATOM	224 CZ2 TRP 185	
	ATOM	225 CZ3 TRP 185	
	ATOM	226 CH2 TRP 185	· · · · · · · · · · · · · · ·
	ATOM	227 C TRP 185	
	ATOM	228 O TRP 185	
25	ATOM	229 N LYS 186	14.277 8.032 5.232 1.00 43.72
	ATOM	230 CA LYS 186	15.694 8.329 5.035 1.00 43.72
	ATOM	231 CB LYS 186	16.353 7.168 4.282 1.00 64.14
	ATOM	232 CG LYS 186	17.830 7.355 3.945 1.00 64.14
	ATOM	233 CD LYS 186	18.758 7.175 5.139 1.00 64.14
30	ATOM	234 CE LYS 186	20.195 7.060 4.652 1.00 64.14
	ATOM	235 NZ LYS 186	20.348 5.838 3.805 1.00 64.14
	ATOM	236 C LYS 186	15.900 9.634 4.263 1.00 43.72
	ATOM	237 O LYS 186	16.948 10.256 4.366 1.00 64.14
	ATOM	238 N GLN 187	14.892 10.032 3.491 1.00 58.06
35	ATOM	239 CA GLN 187	14.958 11.244 2.682 1.00 58.06
	ATOM	240 CB GLN 187	14.288 10.997 1.321 1.00 74.68
	ATOM	241 CG GLN 187	14.639 9.662 0.667 1.00 74.68
	ATOM	242 CD GLN 187	16.133 9.397 0.607 1.00 74.68
	ATOM	243 OE1 GLN 187	16.926 10.312 0.381 1.00 74.68
40	ATOM	244 NE2 GLN 187	
	ATOM	245 C GLN 187	14.322 12.466 3.342 1.00 58.06
	ATOM	246 O GLN 187	14.897 13.551 3.358 1.00 74.68
	ATOM	247 N ARG 188	13.117 12.280 3.866 1.00 54.11
	ATOM	248 CA ARG 188	12.363 13.360 4.505 1.00 54.11
45	ATOM	249 CB ARG 188	10.889 13.115 4.334 1.00 53.33
-	ATOM	250 C ARG 188	12.654 13.626 5.977 1.00 54.11
	ATOM	251 O ARG 188	11.879 14.298 6.659 1.00 53.33
	ATOM	252 N ARG 189	13.754 13.090 6.473 1.00 39.52
	ATOM	253 CA ARG 189	
50	ATOM	254 CB ARG 189	0 100 1 00 50 05
		-	

	ATOM	255 CG ARG 189	15.969 11.555 7.991 1.00 60.85
	ATOM	256 CD ARG 189	16.442 10.298 8.693 1.00 60.85
	ATOM	257 NE ARG 189	17.833 9.963 8.385 1.00 60.85
	ATOM	258 CZ ARG 189	18.627 9.261 9.190 1.00 60.85
5	ATOM	259 NH1 ARG 189	18.178 8.805 10.356 1.00 60.85
,	ATOM	260 NH2 ARG 189	19.882 9.021 8.841 1.00 60.85
	ATOM	261 C ARG 189	15.109 14.378 8.109 1.00 39.52
	ATOM	262 O ARG 189	16.037 14.565 7.320 1.00 60.85
	ATOM	263 N LYS 190	14.934 15.100 9.212 1.00 44.13
10	ATOM	264 CA LYS 190	15.834 16.183 9.586 1.00 44.13
10	ATOM	265 CB LYS 190	15.068 17.500 9.680 1.00 45.33
	ATOM	266 C LYS 190	16.472 15.846 10.928 1.00 44.13
	ATOM	267 O LYS 190	15.827 15.272 11.805 1.00 45.33
	ATOM	268 N PHE 191	17.748 16.184 11.067 1.00 35.64
1.5		269 CA PHE 191	18.489 15.928 12.291 1.00 35.64
15	ATOM	270 CB PHE 191	19.993 16.008 12.025 1.00 53.94
	ATOM		20.550 14.827 11.286 1.00 53.94
	ATOM		20.209 14.596 9.958 1.00 53.94
	ATOM	272 CD1 PHE 191	21.430 13.949 11.915 1.00 53.94
	ATOM	273 CD2 PHE 191	20.735 13.510 9.265 1.00 53.94
20	ATOM	274 CE1 PHE 191	21.964 12.859 11.230 1.00 53.94
	ATOM	275 CE2 PHE 191	21.964 12.839 11.230 1.00 33.94 21.615 12.639 9.900 1.00 53.94
	ATOM	276 CZ PHE 191	18.135 16.928 13.384 1.00 35.64
	ATOM	277 C PHE 191	17.997 18.127 13.120 1.00 53.94
~ ~	ATOM	278 O PHE 191	
25	ATOM	279 N LEU 192	17.978 16.439 14.610 1.00 44.53 17.683 17.315 15.736 1.00 44.53
	ATOM	280 CA LEU 192	17.326 16.493 16.980 1.00 22.94
	ATOM	281 CB LEU 192	
	ATOM	282 CG LEU 192	16.931 17.259 18.246 1.00 22.94 15.568 17.906 18.064 1.00 22.94
•	ATOM	283 CD1 LEU 192	16.909 16.308 19.427 1.00 22.94
30	ATOM	284 CD2 LEU 192	
	ATOM	285 C LEU 192	
	ATOM	286 O LEU 192	20.049 17.507 16.129 1.00 22.94 18.895 19.444 15.977 1.00 34.26
	ATOM	287 N PRO 193	
	ATOM	288 CD PRO 193	17.670 20.241 15.781 1.00 46.23
35	ATOM	289 CA PRO 193	20.058 20.311 16.198 1.00 34.26 19.417 21.670 16.465 1.00 46.23
	ATOM	290 CB PRO 193	
	ATOM	291 CG PRO 193	18.213 21.641 15.579 1.00 46.23
	ATOM	292 C PRO 193	20.917 19.844 17.372 1.00 34.26
	ATOM	293 O PRO 193	20.413 19.614 18.471 1.00 46.23
40	ATOM	294 N ASP 194	22.217 19.716 17.125 1.00 42.67
	ATOM	295 CA ASP 194	23.174 19.254 18.128 1.00 42.67
	ATOM	296 CB ASP 194	24.583 19.226 17.536 1.00 68.50
	ATOM	297 CG ASP 194	24.731 18.185 16.450 1.00 68.50
	ATOM	298 OD1 ASP 194	25.066 17.027 16.782 1.00 68.50
45	ATOM	299 OD2 ASP 194	24.498 18.518 15.269 1.00 68.50
	ATOM	300 C ASP 194	23.187 20.003 19.457 1.00 42.67
	ATOM	301 O ASP 194	23.545 19.432 20.486 1.00 68.50
	ATOM	302 N ASP 195	22.817 21.280 19.438 1.00 47.52
	ATOM	303 CA ASP 195	22.793 22.070 20.666 1.00 47.52
50	ATOM	304 CB ASP 195	22.586 23.559 20.351 1.00 85.02

ATOM 306 ODI ASP 195 ATOM 306 ODI ASP 195 ATOM 307 OD2 ASP 195 ATOM 307 OD2 ASP 195 ATOM 308 C ASP 195 ATOM 308 C ASP 195 ATOM 309 O ASP 195 ATOM 310 N ILE 196 ATOM 311 CA ILE 196 ATOM 312 CB ILE 196 ATOM 315 CGI ILE 196 ATOM 316 C ILE 196 ATOM 317 O ILE 196 ATOM 317 O ILE 196 ATOM 318 N GLY 197 ATOM 319 CA GLY 197 ATOM 320 C GLY 197 ATOM 321 O GLY 197 ATOM 322 N GLN 198 ATOM 323 CA GLN 198 ATOM 325 C GLN 198 ATOM 326 C GLN 198 ATOM 327 N SER 199 ATOM 327 N SER 199 ATOM 330 OG SER 199 ATOM 331 OG SER 199 ATOM 331 CA SER 199 ATOM 332 O SER 199 ATOM 333 N PRO 200 ATOM 334 CB PRO 200 ATOM 335 CA PRO 200 ATOM 336 CB PRO 200 ATOM 336 CB PRO 200 ATOM 337 CG ILE 201 ATOM 338 CA PRO 200 ATOM 330 OG SER 199 ATOM 331 C BILE 201 ATOM 332 CB ILE 201 ATOM 334 CB PRO 200 ATOM 335 CB PRO 200 ATOM 341 CGI ILE 201 ATOM 341 CGI ILE 201 ATOM 342 CB ILE 201 ATOM 343 CGI ILE 201 ATOM 344 CGI ILE 201 ATOM 345 CB VAL 202 ATOM 345 CB VAL 202 ATOM 346 C ILE 201 ATOM 347 O ILE 201 ATOM 348 N VAL 202 ATOM 348 N VAL 202 ATOM 355 CGI VAL 202 ATOM 355 CGI VAL 202 ATOM 356 CO VAL 202 ATOM 357 CGI VAL 202 ATOM 367 CGI VAL 202 ATOM 376 CB VAL 202 ATOM 377 CGI VAL 202 ATOM 377 CGI VAL 202 ATOM 378 CRO VAL 202 ATOM 379 CRO VAL 202 ATOM 350 CGI VAL 202 ATOM 350 CGI VAL 202 ATOM 351 CGI VAL 202 ATOM 351 CGI VAL 202 ATOM 352 CGI VAL 202 ATOM 353 C VAL 202 ATOM 353 C VAL 202 ATOM 354 C VAL 202 ATOM 355 CGI VAL 202 ATOM 356 CRO VAL 202 ATOM 357 CGI VAL 202 ATOM 357 CGI VAL 202 ATOM 358 CRO VAL 202 ATOM 357 CGI VAL 202 ATOM 357 CGI VAL 202 ATOM 357 CGI VAL 202 ATOM 358 CRO VAL 202 ATOM 357 CGI VAL 202 ATOM 358 CRO VAL		ATON	205 CC ACD 105	21 227 22 824 10 527 1 00 85 02
ATOM 307 OD2 ASP 195 21.377 23.683 18.294 1.00 85.02 ATOM 308 C ASP 195 21.715 21.561 21.627 1.00 47.52 ATOM 309 O ASP 195 21.715 21.561 21.627 1.00 47.52 ATOM 310 N ILE 196 20.760 20.810 21.089 1.00 44.54 ATOM 311 CA ILE 196 19.663 20.259 21.875 1.00 44.54 ATOM 312 CB ILE 196 18.379 20.137 21.023 1.00 39.66 ATOM 315 CD1 ILE 196 18.031 21.496 20.407 1.00 39.66 ATOM 316 C ILE 196 20.300 18.882 22.420 1.00 44.54 ATOM 317 O ILE 196 ATOM 318 N GLY 197 20.301 18.882 22.420 1.00 44.54 ATOM 319 CA GLY 197 20.006 17.372 24.307 1.00 39.66 ATOM 320 C GLY 197 20.006 17.372 24.307 1.00 42.85 ATOM 321 O GLY 197 21.371 17.285 24.956 1.00 42.85 ATOM 322 N GLN 198 23.351 18.444 25.754 1.00 53.07 ATOM 323 CA GLN 198 23.351 18.444 25.754 1.00 53.07 ATOM 326 O GLN 198 24.357 19.103 24.810 1.00 44.23 ATOM 326 O GLN 198 24.357 19.103 24.810 1.00 44.23 ATOM 320 C GLY 197 21.171 1.00 53.07 ATOM 328 CA SER 199 22.079 19.18 29.019 1.00 35.30 ATOM 320 C GLN 198 24.396 19.545 27.616 1.00 44.23 ATOM 330 OG SER 199 22.079 19.18 29.019 1.00 35.30 ATOM 331 C SER 199 21.472 21.328 28.806 1.00 58.72 ATOM 333 N PRO 200 21.544 17.928 30.387 1.00 34.70 ATOM 336 CB PRO 200 20.740 17.184 31.362 1.00 34.70 ATOM 337 CG PRO 200 20.740 17.184 31.362 1.00 34.70 ATOM 340 N ILE 201 ATOM 341 CA ILE 201 ATOM 342 CB ILE 201 ATOM 342 CB ILE 201 ATOM 345 CD IILE 201 ATOM 346 C ILE 201 ATOM 347 O ILE 201 ATOM 347 O ILE 201 ATOM 348 N VAL 202 14.688 20.991 34.078 1.00 51.32 ATOM 350 CB VAL 202 16.661 17.787 36.043 1.00 50.33 ATOM 350 CB VAL 202 16.661 17.787 36.043 1.00 50.33 ATOM 350 CB VAL 202 16.661 17.787 36.043 1.00 36.59 ATOM 350 CB VAL 202 16.661 17.787 36.043 1.00 50.33 ATOM 350 CB VAL 202 14.698 19.074 34.714 1.00 36.59 ATOM 351 CGI VAL 202 14.698 19.074 34.714 1.00 36.59 ATOM 353 C VAL 202 14.698 19.074 34.714 1.00 36.59 ATOM 353 C VAL 202 14.698 19.074 34.714 1.00 36.59 ATOM 353 C VAL 202 14.698 19.074 34.714 1.00 36.59 ATOM 353 C VAL 202 14.698 19.074 34.714 1.00 36.59 ATOM 353 C VAL 202 14.202 17.311 36.304 1.00 36.59 ATOM 353 C VAL 202 14.202 17.31				
ATOM 308 C ASP 195 ATOM 309 O ASP 195 ATOM 310 N ILE 196 ATOM 311 CA ILE 196 ATOM 311 CA ILE 196 ATOM 312 CB ILE 196 ATOM 313 CG2 ILE 196 ATOM 315 CD1 ILE 196 ATOM 316 C ILE 196 ATOM 317 O ILE 196 ATOM 319 CA GLY 197 ATOM 320 C GLY 197 ATOM 321 C GILY 197 ATOM 322 N GLN 198 ATOM 322 N GLN 198 ATOM 323 CA GLN 198 ATOM 325 C GLN 198 ATOM 326 O GLN 198 ATOM 327 N SER 199 ATOM 327 N SER 199 ATOM 329 CB SER 199 ATOM 330 C SER 199 ATOM 331 C SER 199 ATOM 331 C SER 199 ATOM 332 C GRO 20 20 20 21.311 15.769 31.20 34.70 ATOM 335 CA PRO 200 ATOM 336 C B PRO 200 ATOM 337 CG PRO 200 ATOM 338 N CAL 202 ATOM 349 CA VAL 202 ATOM 349 CA VAL 202 ATOM 340 N 1.00 36.59 ATOM 350 CB VAL 202 ATOM 340 N 1.00 36.59 ATOM 340 N ILE 201 ATOM 340 N ILE 201 ATOM 340 CA VAL 202 ATOM 340 CA VAL 202 ATOM 340 N 36 CB VAL 202 ATOM 351 C VAL 202 ATOM 353 C VAL 202 ATOM 355 CAGLN 100 35.30 ATOM 360 CB VAL 202 ATOM 375 CAGLN 100 35.30 ATOM 376 CB VAL 202 ATOM 377 N 358 CG SCA				
5         ATOM         309         O         ASP         195         21.762         21.826         22.831         1.00         84.502           ATOM         310         N         ILE         196         20.760         20.810         21.089         1.00         44.54           ATOM         312         CB         ILE         196         18.379         20.137         21.023         1.00         39.66           ATOM         314         CGI         ILE         196         17.223         19.627         21.874         1.00         39.66           ATOM         315         CDI         ILE         196         17.223         19.627         21.874         1.00         39.66           ATOM         316         C         ILE         196         16.816         21.475         19.503         1.00         39.66           ATOM         317         O         ILE         196         20.30         18.882         22.420         1.00         44.54           ATOM         319         CA         GLY         197         20.006         17.372         24.307         1.00         42.85           ATOM         321         O         GLY		_		· ·
ATOM 310 N ILE 196 ATOM 311 CA ILE 196 ATOM 312 CB ILE 196 ATOM 313 CG2 ILE 196 ATOM 313 CG2 ILE 196 ATOM 314 CG1 ILE 196 ATOM 315 CD1 ILE 196 ATOM 315 CD1 ILE 196 ATOM 316 C ILE 196 ATOM 317 O ILE 196 ATOM 317 O ILE 196 ATOM 318 N GLY 197 ATOM 320 C GLY 197 ATOM 320 C GLY 197 ATOM 321 O GLY 197 ATOM 322 N GLN 198 ATOM 323 CA GLN 198 ATOM 324 CB GLN 198 ATOM 325 C GLN 198 ATOM 326 O GLN 198 ATOM 327 N SER 199 ATOM 328 CA SER 199 ATOM 330 OG SER 199 ATOM 330 OG SER 199 ATOM 331 C SER 199 ATOM 331 C SER 199 ATOM 332 O SER 199 ATOM 333 N PRO 200 ATOM 334 CD PRO 200 ATOM 335 CA PRO 200 ATOM 336 CB PRO 200 ATOM 337 CG PRO 200 ATOM 344 CB ILE 201 ATOM 344 CB ILE 201 ATOM 345 CB VAL 202 ATOM 346 C ILE 201 ATOM 347 O ILE 201 ATOM 348 N VAL 202 ATOM 349 CA VAL 202 ATOM 350 CB VAL 202 ATOM 349 CA VAL 202 ATOM 340 CB ILA 204 ATOM 340 CB ILE 201 ATOM 340 CB ILE 201 ATOM 346 CB ILE 201 ATOM 347 O ILE 201 ATOM 348 N VAL 202 ATOM 350 CB VAL 202 ATOM 351 CG IVAL 202 ATOM 355 CA VAL 202 ATOM 360 CB VAL 202 A	_			
ATOM 311 CA ILE 196	5			
ATOM 312 CB ILE 196 18.379 20.137 21.023 1.00 39.66 ATOM 313 CG2 ILE 196 17.223 19.627 21.874 1.00 39.66 ATOM 314 CGI ILE 196 16.816 21.475 19.503 1.00 39.66 ATOM 315 CD1 ILE 196 16.816 21.475 19.503 1.00 39.66 ATOM 316 C ILE 196 20.030 18.882 22.420 1.00 44.54 ATOM 317 O ILE 196 20.582 18.046 21.705 1.00 39.66 ATOM 318 N GLY 197 20.006 17.372 24.307 1.00 42.85 ATOM 320 C GLY 197 20.006 17.372 24.307 1.00 42.85 ATOM 321 O GLY 197 21.815 16.198 25.318 1.00 42.285 ATOM 322 N GLN 198 22.029 18.425 25.137 1.00 53.07 ATOM 323 CA GLN 198 23.351 18.444 25.754 1.00 53.07 ATOM 325 C GLN 198 24.357 19.103 24.810 1.00 44.23 ATOM 326 O GLN 198 24.357 19.103 24.810 1.00 44.23 ATOM 327 N SER 199 22.170 19.244 27.729 1.00 35.30 ATOM 329 CB SER 199 22.037 19.918 29.019 1.00 35.30 ATOM 331 C SER 199 22.037 19.918 29.019 1.00 35.30 ATOM 332 O SER 199 22.037 19.918 29.019 1.00 35.30 ATOM 331 C SER 199 22.037 19.918 29.019 1.00 35.30 ATOM 332 O SER 199 22.037 19.918 29.019 1.00 35.30 ATOM 333 N PRO 200 21.544 17.928 30.387 1.00 44.70 ATOM 335 CA PRO 200 20.740 17.184 31.362 1.00 38.71 ATOM 336 CB PRO 200 20.740 17.184 31.362 1.00 38.71 ATOM 337 CG PRO 200 22.737 15.992 30.878 1.00 34.70 ATOM 340 N ILE 201 19.961 19.041 34.604 1.00 42.94 ATOM 340 CB ILE 201 19.069 17.784 32.759 1.00 34.70 ATOM 340 CB ILE 201 19.069 17.784 32.759 1.00 34.70 ATOM 340 N ILE 201 19.961 19.041 34.604 1.00 42.94 ATOM 344 CGI ILE 201 19.069 21.511 33.510 1.00 51.32 ATOM 346 C ILE 201 19.069 11.13 35.10 1.00 51.32 ATOM 346 C ILE 201 19.069 11.13 35.10 1.00 51.32 ATOM 346 C ILE 201 19.069 11.13 35.10 1.00 51.32 ATOM 347 O ILE 201 19.069 11.13 35.10 1.00 51.32 ATOM 348 N VAL 202 17.377 18.172 35.336 1.00 36.59 ATOM 350 C B VAL 202 15.296 17.722 35.326 1.00 36.59 ATOM 350 C B VAL 202 16.661 17.787 36.043 1.00 50.33 ATOM 350 C B VAL 202 15.296 17.722 35.326 1.00 36.59 ATOM 353 C VAL 202 14.202 17.007 16.435 36.665 1.00 36.59 ATOM 353 C VAL 202 14.202 17.007 16.435 36.665 1.00 36.33				
ATOM 314 CG1 ILE 196				19.663 20.259 21.875 1.00 44.54
10 ATOM 314 CG1 ILE 196 ATOM 315 CD1 ILE 196 ATOM 315 CD1 ILE 196 ATOM 316 C ILE 196 ATOM 316 C ILE 196 ATOM 317 O ILE 196 ATOM 318 N GLY 197 20.003 18.882 22.420 1.00 44.54 ATOM 318 N GLY 197 20.006 17.372 24.307 1.00 39.66 ATOM 318 N GLY 197 20.006 17.372 24.307 1.00 42.85 ATOM 320 C GLY 197 21.371 17.285 24.956 1.00 42.85 ATOM 321 O GLY 197 21.815 16.198 25.318 1.00 40.22 ATOM 322 N GLN 198 22.029 18.425 25.137 1.00 53.07 ATOM 323 CA GLN 198 24.357 19.103 24.810 1.00 44.23 ATOM 325 C GLN 198 24.357 19.103 24.810 1.00 44.23 ATOM 325 C GLN 198 24.356 19.545 27.616 1.00 44.23 ATOM 329 CB SER 199 22.170 19.244 27.729 1.00 35.30 ATOM 330 OG SER 199 22.037 19.918 29.019 1.00 35.30 ATOM 331 C SER 199 22.037 19.918 29.019 1.00 58.72 ATOM 331 C SER 199 21.472 21.328 28.806 1.00 58.72 ATOM 333 N PRO 200 21.541 17.928 30.387 1.00 34.70 ATOM 336 CB PRO 200 ATOM 336 CB PRO 200 ATOM 337 CG PRO 200 ATOM 338 C PRO 200 20.740 17.184 31.362 1.00 34.70 ATOM 340 N ILE 201 ATOM 340 N ILE 201 ATOM 341 CA ILE 201 ATOM 344 CG1 ILE 201 ATOM 345 CD1 ILE 201 ATOM 346 C ILE 201 ATOM 347 O ILE 201 ATOM 348 N VAL 202 ATOM 351 CG1 VAL 202 ATOM 352 CG2 VAL 202 ATOM 352 CG2 VAL 202 ATOM 353 C VAL 202 ATOM 353 C COL ATOM 354 CD PRO 200 ATOM 348 N VAL 202 ATOM 355 CG6 VAL 202 ATOM 350 CB VAL 202 ATOM 350 CB VAL 202 ATOM 351 CG1 VAL 202 ATOM 352 CG2 VAL 202 ATOM 353 C VAL 202 ATOM 350 CB VAL 202 ATOM 351 CG1 VAL 202 ATOM 353 C VAL 202 ATO				
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35       ATOM       339       O       PRO       200       22.006       17.692       33.341       1.00       38.71         ATOM       340       N       ILE       201       19.876       18.413       33.286       1.00       42.94         ATOM       341       CA       ILE       201       19.961       19.041       34.604       1.00       42.94         ATOM       342       CB       ILE       201       20.059       20.582       34.491       1.00       51.32         ATOM       343       CG2       ILE       201       21.468       20.991       34.078       1.00       51.32         ATOM       344       CG1       ILE       201       19.099       21.111       33.510       1.00       51.32         ATOM       345       CD1       ILE       201       19.169       22.582       33.164       1.00       51.32         ATOM       346       C       ILE       201       18.871       18.676       35.610       1.00       42.94         ATOM       347       O       ILE       201       19.049       18.875       36.814       1.00       51.32         ATOM				
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40       ATOM       344       CG1 ILE       201       19.009       21.111       33.510       1.00 51.32         ATOM       345       CD1 ILE       201       19.169       22.582       33.164       1.00 51.32         ATOM       346       C       ILE       201       18.871       18.676       35.610       1.00 42.94         ATOM       347       O       ILE       201       19.049       18.875       36.814       1.00 51.32         ATOM       348       N       VAL       202       17.737       18.172       35.133       1.00 50.33         ATOM       349       CA       VAL       202       16.661       17.787       36.043       1.00 50.33         ATOM       350       CB       VAL       202       15.296       17.722       35.326       1.00 36.59         ATOM       351       CG1       VAL       202       14.202       17.311       36.304       1.00 36.59         ATOM       352       CG2       VAL       202       14.968       19.074       34.714       1.00 36.59         ATOM       353       C       VAL       202       17.007       16.435       36.665       1.00 50.33 </td <td></td> <td></td> <td></td> <td></td>				
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ATOM 347 O ILE 201 19.049 18.875 36.814 1.00 51.32 ATOM 348 N VAL 202 17.737 18.172 35.133 1.00 50.33 45 ATOM 349 CA VAL 202 16.661 17.787 36.043 1.00 50.33 ATOM 350 CB VAL 202 15.296 17.722 35.326 1.00 36.59 ATOM 351 CG1 VAL 202 14.202 17.311 36.304 1.00 36.59 ATOM 352 CG2 VAL 202 14.968 19.074 34.714 1.00 36.59 ATOM 353 C VAL 202 17.007 16.435 36.665 1.00 50.33				
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ATOM 353 C VAL 202 17.007 16.435 36.665 1.00 50.33		<b>ATOM</b>	351 CG1 VAL 202	
		<b>ATOM</b>	352 CG2 VAL 202	
		ATOM	353 C VAL 202	
	50	ATOM	354 O VAL 202	17.335 15.481 35.955 1.00 36.59

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16.960 16.375 37.991 1.00 49.46
            355 N SER 203
    ATOM
                               17.289 15.166 38.736 1.00 49.46
            356 CA SER 203
    ATOM
                               17.298 15.467 40.241 1.00 64.20
            357 CB SER 203
    ATOM
                               17.673 14.330 41.003 1.00 64.20
            358 OG SER 203
    ATOM
                               16.356 13.992 38.463 1.00 49.46
            359 C SER 203
    ATOM
                               15.147 14.166 38.310 1.00 64.20
            360 O SER 203
    ATOM
                               16.944 12.800 38.419 1.00 41.99
            361 N MET 204
    ATOM
                                16.223 11.551 38.205 1.00 41.99
            362 CA MET 204
    ATOM
            363 CB MET 204
                                16.320 11.096 36.746 1.00 48.64
    ATOM
                                15.470 11.895 35.771 1.00 48.64
            364 CG MET 204
10
    ATOM
                                13.702 11.783 36.114 1.00 48.64
    ATOM
            365 SD MET 204
                                13.284 10.257 35.264 1.00 48.64
            366 CE MET 204
    ATOM
                               16.900 10.528 39.109 1.00 41.99
    ATOM
            367 C MET 204
            368 O MET 204
369 N PRO 205
                               18.127 10.417 39.121 1.00 48.64
    ATOM
                               16.108 9.754 39.869 1.00 38.42
    ATOM
15
                               14.633 9.815 39.866 1.00 52.20
    ATOM
            370 CD PRO 205
                                16.586 8.724 40.797 1.00 38.42
            371 CA PRO 205
    ATOM
                                15.334 7.888 41.041 1.00 52.20
            372 CB PRO 205
    ATOM
                               14.254 8.919 41.028 1.00 52.20
           373 CG PRO 205
    ATOM
    ATOM 374 C PRO 205
                               17.769 7.858 40.340 1.00 38.42
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    ATOM 375 O PRO 205
                               18.724 7.675 41.092 1.00 52.20
            376 N ASP 206
                               17.720 7.349 39.111 1.00 49.06
    ATOM
    ATOM
            377 CA ASP 206
                               18.791 6.490 38.601 1.00 49.06
                               18.282 5.627 37.437 1.00 74.42
            378 CB ASP 206
    ATOM
                                17.690 6.450 36.305 1.00 74.42
            379 CG ASP 206
25
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                                18.397 7.335 35.770 1.00 74.42
            380 OD1 ASP 206
    ATOM
            381 OD2 ASP 206
                                16.516 6.199 35.948 1.00 74.42
    ATOM
            382 C ASP 206
383 O ASP 206
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     ATOM
                               21.069 6.506 37.838 1.00 74.42
     ATOM
            384 N GLY 207
                               20.139 8.505 38.272 1.00 42.48
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    ATOM
                               21.355 9.225 37.928 1.00 42.48
            385 CA GLY 207
     ATOM
                               21.330 9.965 36.601 1.00 42.48
            386 C GLY 207
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     ATOM 387 O GLY 207
                               21.890 11.058 36.494 1.00 42.50
     ATOM 388 N ASP 208
                               20.725 9.365 35.581 1.00 46.70
                                20.636 9.999 34.266 1.00 46.70
            389 CA ASP 208
35
     ATOM
                                20.162 8.994 33.212 1.00 61.56
            390 CB ASP 208
     ATOM
                                21.143 7.856 33.006 1.00 61.56
            391 CG ASP 208
     ATOM
                                20.723 6.684 33.122 1.00 61.56
            392 OD1 ASP 208
     ATOM
            393 OD2 ASP 208
                                22.330 8.134 32.724 1.00 61.56
     ATOM
            394 C ASP 208
                               19.666 11.176 34.339 1.00 46.70
40
    ATOM
            395 O ASP 208
396 N LYS 209
                               18.462 10.983 34.506 1.00 61.56
     ATOM
                               20,200 12.389 34.238 1.00 41.30
     ATOM
                                19.389 13.602 34.308 1.00 41.30
            397 CA LYS 209
     ATOM
                                20.254 14.782 34.732 1.00 41.38
            398 CB LYS 209
     ATOM
                               18.657 13.916 33.004 1.00 41.30
            399 C LYS 209
45
     ATOM
                               19.052 13.458 31.930 1.00 41.38
            400 O LYS 209
     ATOM
                               17.603 14.723 33.109 1.00 43.36
            401 N VAL 210
     ATOM
            402 CA VAL 210
403 CB VAL 210
                                16.792 15.107 31.954 1.00 43.36
     ATOM
                                15.275 15.014 32.282 1.00 30.23
     ATOM
            404 CG1 VAL 210
                                14.440 15.358 31.055 1.00 30.23
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     ATOM
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	ATOM	405 CG2 VAL 210	14.923 13.624 32.782 1.00 30.23
	ATOM	406 C VAL 210	17.088 16.522 31.442 1.00 43.36
	ATOM	407 O VAL 210	17.395 17.430 32.221 1.00 30.23
	ATOM	408 N ASP 211	17.004 16.685 30.125 1.00 27.49
5	ATOM	409 CA ASP 211	17.217 17.966 29.458 1.00 27.49
	ATOM	410 CB ASP 211	18.073 17.765 28.198 1.00 30.75
	ATOM	411 CG ASP 211	18.360 19.068 27.447 1.00 30.75
	ATOM	412 OD1 ASP 211	19.473 19.196 26.900 1.00 30.75
	ATOM	413 OD2 ASP 211	17.484 19.955 27.370 1.00 30.75
10	ATOM	414 C ASP 211	15.819 18.445 29.073 1.00 27.49
	ATOM	415 O ASP 211	15.197 17.892 28.166 1.00 30.75
	ATOM	416 N LEU 212	15.343 19.488 29.745 1.00 31.99
	ATOM	417 CA LEU 212	14.013 20.042 29.492 1.00 31.99
	ATOM	418 CB LEU 212	13.778 21.274 30.369 1.00 35.19
15	ATOM	419 CG LEU 212	13.606 20.997 31.864 1.00 35.19
	ATOM	420 CD1 LEU 212	13.621 22.298 32.652 1.00 35.19
	ATOM	421 CD2 LEU 212	12.309 20.237 32.098 1.00 35.19
	ATOM	422 C LEU 212	13.713 20.377 28.032 1.00 31.99
	ATOM	423 O LEU 212	12.625 20.083 27.539 1.00 35.19
20	ATOM	424 N GLU 213	14.672 20.981 27.338 1.00 28.70
	ATOM	425 CA GLU 213	14.468 21.345 25.940 1.00 28.70
	ATOM	426 CB GLU 213	15.623 22.209 25.428 1.00 62.21
	ATOM	427 CG GLU 213	15.434 22.707 23.997 1.00 62.21
	ATOM	428 CD GLU 213	16.651 23.440 23.446 1.00 62.21
25	ATOM	429 OE1 GLU 213	17.778 23.214 23.945 1.00 62.21
	ATOM	430 OE2 GLU 213	16.478 24.237 22.498 1.00 62.21
	ATOM	431 C GLU 213	14.317 20.104 25.067 1.00 28.70
	ATOM	432 O GLU 213	13.403 20.024 24.247 1.00 62.21
	ATOM	433 N ALA 214	15.201 19.130 25.262 1.00 28.17
30	ATOM	434 CA ALA 214	15.162 17.890 24.494 1.00 28.17
	ATOM	435 CB ALA 214	16.330 16.998 24.872 1.00 42.74
	ATOM	436 C ALA 214	13.844 17.176 24.759 1.00 28.17
	ATOM	437 O ALA 214	13.174 16.726 23.829 1.00 42.74
	ATOM	438 N PHE 215	13.468 17.104 26.032 1.00 21.66
35	ATOM	439 CA PHE 215	12,222 16.471 26.444 1.00 21.66
	ATOM	440 CB PHE 215	12.033 16.628 27.958 1.00 28.76
	ATOM	441 CG PHE 215	10.751 16.038 28.481 1.00 28.76
	ATOM	442 CD1 PHE 215	10.675 14.689 28.815 1.00 28.76
	ATOM	443 CD2 PHE 215	9.623 16.835 28.653 1.00 28.76
40	ATOM	444 CE1 PHE 215	9.493 14.143 29.315 1.00 28.76
	ATOM	445 CE2 PHE 215	8.438 16.300 29.150 1.00 28.76
	ATOM	446 CZ PHE 215	8.373 14.951 29.482 1.00 28.76
	ATOM	447 C PHE 215	11.068 17.132 25.696 1.00 21.66
	ATOM	448 O PHE 215	10.215 16.451 25.122 1.00 28.76
45	ATOM	449 N SER 216	11.073 18.462 25.680 1.00 28.03
	ATOM	450 CA SER 216	10.043 19.242 25.007 1.00 28.03
	ATOM	451 CB SER 216	10.349 20.734 25.146 1.00 33.85
	ATOM	452 OG SER 216	9.300 21.529 24.624 1.00 33.85
	ATOM	453 C SER 216	9.945 18.857 23.532 1.00 28.03
50	ATOM	454 O SER 216	8.852 18.613 23.019 1.00 33.85
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    ATOM
             457 CB GLU 217
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    ATOM
                                13.174 19.815 20.811 1.00 47.68
             458 CG GLU 217
     ATOM
                                12.405 20.684 19.829 1.00 47.68
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    ATOM
                                 12.542 20.465 18.606 1.00 47.68
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    ATOM
            463 O GLU 217
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     ATOM
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             467 CG PHE 218
             468 CD1 PHE 218
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     ATOM
                                12.932 13.677 21.144 1.00 24.12
             469 CD2 PHE 218
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                                14.793 13.484 23.187 1.00 24.12
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     ATOM
             471 CE2 PHE 218
                                14.290 13.607 20.843 1.00 24.12
     ATOM
                                15.221 13.511 21.867 1.00 24.12
            472 CZ PHE 218
     ATOM
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     ATOM
            473 C PHE 218
                                7.985 14.166 21.369 1.00 24.12
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     ATOM
            474 O PHE 218
                                8.358 15.227 23.312 1.00 20.07
            475 N THR 219
     ATOM
            476 CA THR 219
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                                 6.741 15.766 25.118 1.00 28.98
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     ATOM
            478 OG1 THR 219
                                 7.275 14.767 26.132 1.00 28.98
            479 CG2 THR 219
25
     ATOM
                                6.080 16.011 22.696 1.00 20.07
             480 C THR 219
     ATOM
                                4.914 15.670 22.482 1.00 28.98
             481 O THR 219
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             482 N LYS 220
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             484 CB LYS 220
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             485 C LYS 220
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             487 N ILE 221
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             488 CA ILE 221
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            489 CB ILE 221
                                6.842 14.915 17.413 1.00 25.19
     ATOM
35
            490 CG2 ILE 221
491 CG1 ILE 221
                                 7.240 16.264 16.838 1.00 25.19
     ATOM
                                 8.050 14.215 18.043 1.00 25.19
     ATOM
                                 9.113 13.799 17.044 1.00 25.19
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             492 CD1 ILE 221
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             494 O ILE 221
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             495 N ILE 222
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     ATOM
                                4.720 12.162 20.687 1.00 24.41
             496 CA ILE 222
     ATOM
             497 CB ILE 222
                                5.189 11.916 22.147 1.00 27.10
     ATOM
                                4.221 12.545 23.145 1.00 27.10
             498 CG2 ILE 222
     ATOM
                                 5.302 10.410 22.400 1.00 27.10
     ATOM
             499 CG1 ILE 222
45
                                 6.062 10.053 23.646 1.00 27.10
             500 CD1 ILE 222
     ATOM
                               3.231 11.845 20.541 1.00 24.41
             501 C ILE 222
     ATOM
                               2.864 10.691 20.307 1.00 27.10
             502 O ILE 222
     ATOM
                                2.378 12.861 20.642 1.00 33.16
             503 N THR 223
     ATOM
                                 0.936 12.653 20.520 1.00 33.16
             504 CA THR 223
50
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	ATOM	505 CB THR 223	0.150 13.974 20.721 1.00 36.84
	<b>ATOM</b>	506 OG1 THR 223	0.352 14.442 22.063 1.00 36.84
	<b>ATOM</b>	507 CG2 THR 223	-1.346 13.764 20.484 1.00 36.84
	ATOM	508 C THR 223	0.536 11.954 19.212 1.00 33.16
5	ATOM	509 O THR 223	-0.156 10.932 19.242 1.00 36.84
	<b>ATOM</b>	510 N PRO 224	0.968 12.482 18.048 1.00 18.75
	ATOM	511 CD PRO 224	1.691 13.735 17.770 1.00 26.12
	ATOM	512 CA PRO 224	0.590 11.805 16.802 1.00 18.75
	ATOM	513 CB PRO 224	1.117 12.747 15.715 1.00 26.12
10	ATOM	514 CG PRO 224	2.221 13.497 16.386 1.00 26.12
	ATOM	515 C PRO 224	1.200 10.402 16.701 1.00 18.75
	ATOM	516 O PRO 224	0.606 9.502 16.101 1.00 26.12
	ATOM	517 N ALA 225	2.368 10.213 17.312 1.00 12.19
	ATOM	518 CA ALA 225	3.040 8.916 17.300 1.00 12.19
15	ATOM	519 CB ALA 225	4.415 9.021 17.943 1.00 20.39
	ATOM	520 C ALA 225	2.187 7.881 18.030 1.00 12.19
	ATOM	521 O ALA 225	1.998 6.764 17.545 1.00 20.39
	ATOM	522 N ILE 226	1.645 8.271 19.179 1.00 14.61
	ATOM	523 CA ILE 226	0.798 7.385 19.971 1.00 14.61
20	ATOM	524 CB ILE 226	0.450 8.025 21.332 1.00 16.10
	ATOM	525 CG2 ILE 226	-0.508 7.132 22.108 1.00 16.10
	ATOM	526 CG1 ILE 226	1.729 8.293 22.132 1.00 16.10
	ATOM	527 CD1 ILE 226	1.509 9.113 23.387 1.00 16.10
	ATOM		-0.499 7.094 19.213 1.00 14.61
25	ATOM	· - ·	-0.986 5.961 19.200 1.00 16.10
	ATOM	530 N THR 227	-1.042 8.123 18.569 1.00 15.93
	ATOM	531 CA THR 227	-2.278 7.997 17.800 1.00 15.93
	ATOM	532 CB THR 227	-2.706 9.360 17.207 1.00 22.37
	ATOM	533 OG1 THR 227	-2.890 10.301 18.273 1.00 22.37
30	ATOM	534 CG2 THR 227	-4.014 9.232 16.434 1.00 22.37
	ATOM	535 C THR 227	-2.149 6.964 16.680 1.00 15.93
	ATOM	536 O THR 227	-3.091 6.217 16.402 1.00 22.37
	ATOM	537 N ARG 228	-0.982 6.916 16.045 1.00 14.49
	ATOM	538 CA ARG 228	-0.750 5.956 14.975 1.00 14.49
35	ATOM	539 CB ARG 228	0.602 6.188 14.307 1.00 33.87
	ATOM	540 CG ARG 228	0.701 7.482 13.540 1.00 33.87
	ATOM	541 CD ARG 228	2.053 7.572 12.868 1.00 33.87
	ATOM	542 NE ARG 228	2.510 8.952 12.793 1.00 33.87
	ATOM	543 CZ ARG 228	3.551 9.431 13.469 1.00 33.87
40	ATOM	544 NH1 ARG 228	4.256 8.634 14.270 1.00 33.87
. •	ATOM	545 NH2 ARG 228	
	ATOM	546 C ARG 228	-0.813 4.531 15.516 1.00 14.49
	ATOM	547 O ARG 228	-1.309 3.632 14.839 1.00 33.87
	ATOM	548 N VAL 229	-0.313 4.327 16.735 1.00 14.80
45	ATOM	549 CA VAL 229	-0.333 3.002 17.352 1.00 14.80
	ATOM	550 CB VAL 229	0.456 2.979 18.683 1.00 13.78
	ATOM	551 CG1 VAL 229	0.339 1.612 19.350 1.00 13.78
	ATOM	552 CG2 VAL 229	1.915 3.312 18.430 1.00 13.78
	ATOM	553 C VAL 229	-1.788 2.602 17.591 1.00 14.80
50	ATOM	554 O VAL 229	-2.185 1.465 17.323 1.00 13.78

	ATOM	555 N VAL 230	-2.588 3.561 18.047 1.00 9.33
	ATOM	556 CA VAL 230	-4.005 3.327 18.292 1.00 9.33
	ATOM	557 CB VAL 230	-4.679 4.564 18.909 1.00 16.07
	ATOM	558 CG1 VAL 230	-6.168 4.319 19.076 1.00 16.07
5	ATOM	559 CG2 VAL 230	-4.038 4.896 20.253 1.00 16.07
,	ATOM	560 C VAL 230	-4.700 2.982 16.981 1.00 9.33
	ATOM	561 O VAL 230	-5.504 2.049 16.929 1.00 16.07
			-4.364 3.719 15.922 1.00 12.71
	ATOM		
10	ATOM	563 CA ASP 231	-4.951 3.496 14.603 1.00 12.71 -4.529 4.596 13.624 1.00 27.08
10	ATOM	564 CB ASP 231	
	ATOM	565 CG ASP 231	
	ATOM	566 OD1 ASP 231	-6.144 6.047 14.624 1.00 27.08
	ATOM	567 OD2 ASP 231	-4.370 6.969 13.723 1.00 27.08
	ATOM	568 C ASP 231	-4.570 2.132 14.049 1.00 12.71
15	ATOM	569 O ASP 231	-5.413 1.436 13.483 1.00 27.08
	ATOM	570 N PHE 232	-3.305 1.755 14.215 1.00 14.33
	ATOM	571 CA PHE 232	-2.823 0.461 13.748 1.00 14.33
	ATOM	572 CB PHE 232	-1.351 0.257 14.134 1.00 16.35
	ATOM	573 CG PHE 232	-0.911 -1.184 14.097 1.00 16.35
20	ATOM	574 CD1 PHE 232	-0.789 -1.862 12.887 1.00 16.35
	ATOM	575 CD2 PHE 232	-0.661 -1.879 15.280 1.00 16.35
	<b>ATOM</b>	576 CE1 PHE 232	-0.430 -3.208 12.851 1.00 16.35
	ATOM	577 CE2 PHE 232	-0.302 -3.224 15.255 1.00 16.35
	<b>ATOM</b>	578 CZ PHE 232	-0.187 -3.890 14.038 1.00 16.35
25	ATOM	579 C PHE 232	-3.670 -0.642 14.368 1.00 14.33
	ATOM	580 O PHE 232	-4.226 -1.482 13.661 1.00 16.35
	ATOM	581 N ALA 233	-3.769 -0.619 15.695 1.00 15.30
	ATOM	582 CA ALA 233	-4.537 -1.607 16.444 1.00 15.30
	<b>ATOM</b>	583 CB ALA 233	-4.413 -1.335 17.938 1.00 12.88
30	ATOM	584 C ALA 233	-6.005 -1.609 16.030 1.00 15.30
	ATOM	585 O ALA 233	-6.627 -2.663 15.902 1.00 12.88
	ATOM	586 N LYS 234	-6.542 -0.419 15.795 1.00 25.69
	ATOM	587 CA LYS 234	-7.933 -0.256 15.401 1.00 25.69
	ATOM	588 CB LYS 234	-8.270 1.234 15.318 1.00 45.91
35	ATOM	589 CG LYS 234	-9.574 1.595 15.979 1.00 45.91
	ATOM	590 CD LYS 234	-9.535 1.268 17.463 1.00 45.91
	ATOM	591 CE LYS 234	-10.938 1.047 18.006 1.00 45.91
	ATOM	592 NZ LYS 234	-11.605 -0.106 17.327 1.00 45.91
	ATOM	593 C LYS 234	-8.240 -0.931 14.067 1.00 25.69
40	ATOM	594 O LYS 234	-9.368 -1.368 13.827 1.00 45.91
	ATOM	595 N LYS 235	-7.234 -1.019 13.204 1.00 17.44
	ATOM	596 CA LYS 235	-7.406 -1.627 11.892 1.00 17.44
	ATOM	597 CB LYS 235	-6.459 -0.975 10.884 1.00 26.26
	ATOM	598 CG LYS 235	-6.757 0.499 10.669 1.00 26.26
45	ATOM	599 CD LYS 235	-5.785 1.141 9.706 1.00 26.26
43			-6.154 2.593 9.460 1.00 26.26
	ATOM		-5.231 3.230 8.484 1.00 26.26
	ATOM		-7.258 -3.146 11.875 1.00 17.44
	ATOM	602 C LYS 235	-7.258 -3.146 11.873 1.00 17.44 -7.365 -3.773 10.817 1.00 26.26
50	ATOM	603 O LYS 235	-7.363 -3.773 10.817 1.00 26.20 -7.015 -3.738 13.040 1.00 21.99
50	ATOM	604 N LEU 236	-/.012 -3./36 13.040 1.00 21.99

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-6.880 -5.187 13.144 1.00 21.99
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            606 CB LEU 236
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            607 CG LEU 236
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            608 CD1 LEU 236
    ATOM
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            609 CD2 LEU 236
    ATOM
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    ATOM
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            611 O LEU 236
     ATOM
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            612 N PRO 237
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            613 CD PRO 237
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            615 CB PRO 237
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     ATOM 623 SD MET 238
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626 O MET 238
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             629 CB PHE 239
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             630 CG PHE 239
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             632 CD2 PHE 239
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            633 CE1 PHE 239
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             704 CD1 ILE 248
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WO 99/26966

PCT/US98/25296

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	ATOM	761 CE MET 256	10.127 4.254 24.782 1.00 19.23
	ATOM	762 C MET 256	6.734 0.978 23.246 1.00 13.77
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	ATOM	770 OE2 GLU 257	5.607 -4.361 26.400 1.00 31.54
	ATOM	771 C GLU 257	6.889 -1.254 21.211 1.00 12.57
	ATOM	772 O GLU 257	7.881 -1.452 20.505 1.00 31.54
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	ATOM	777 CG1 ILE 258	3.288 -1.886 19.211 1.00 15.57
	ATOM	778 CD1 ILE 258	1.798 -1.872 18.922 1.00 15.57
25	ATOM	779 C ILE 258	6.289 0.535 18.811 1.00 17.89
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	ATOM	781 N MET 259	6.196 1.636 19.556 1.00 11.23
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••	ATOM	783 CB MET 259	6.568 3.995 20.175 1.00 22.19
30	ATOM	784 CG MET 259	5.112 4.439 20.117 1.00 22.19
	ATOM	785 SD MET 259	4.828 6.033 20.915 1.00 22.19
	ATOM	786 CE MET 259	5.038 5.606 22.621 1.00 22.19
	ATOM	787 C MET 259	8.415 2.637 19.131 1.00 11.23
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35	ATOM	789 N SER 260	8.974 1.994 20.153 1.00 8.59
	ATOM	790 CA SER 260	10.408 1.706 20.195 1.00 8.59
	ATOM	791 CB SER 260	10.763 0.939 21.472 1.00 23.39
	ATOM	792 OG SER 260	10.430 1.685 22.623 1.00 23.39
	ATOM	793 C SER 260	10.793 0.864 18.977 1.00 8.59
40	ATOM	794 O SER 260	11.824 1.100 18.350 1.00 23.39
	ATOM	795 N LEU 261	9.952 -0.111 18.644 1.00 13.26
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	ATOM	800 CD2 LEU 261	7.785 -3.772 16.174 1.00 14.32
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	ATOM	803 N ARG 262	9.330 0.744 16.043 1.00 10.57
50	ATOM	804 CA ARG 262	9.278 1.598 14.861 1.00 10.57

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	ATOM	805 CB ARG 262	8.018 2.454 14.917 1.00 16.08
	<b>ATOM</b>	806 CG ARG 262	6.755 1.647 14.728 1.00 16.08
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	ATOM	808 NE ARG 262	4.418 1.765 14.076 1.00 16.08
5	ATOM	809 CZ ARG 262	3.260 2.289 13.689 1.00 16.08
	ATOM	810 NH1 ARG 262	3.050 3.596 13.780 1.00 16.08
	<b>ATOM</b>	811 NH2 ARG 262	2.322 1.497 13.183 1.00 16.08
	ATOM	812 C ARG 262	10.530 2.471 14.704 1.00 10.57
	ATOM	813 O ARG 262	11.038 2.649 13.589 1.00 16.08
10	ATOM	814 N ALA 263	11.016 3.014 15.820 1.00 13.37
	ATOM	815 CA ALA 263	12.221 3.842 15.831 1.00 13.37
	<b>ATOM</b>	816 CB ALA 263	12.363 4.516 17.172 1.00 17.12
	ATOM	817 C ALA 263	13.443 2.964 15.561 1.00 13.37
	ATOM	818 O ALA 263	14.313 3.316 14.762 1.00 17.12
15	ATOM	819 N ALA 264	13.474 1.802 16.207 1.00 16.55
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	ATOM	822 C ALA 264	14.770 0.364 14.642 1.00 16.55
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20	ATOM	824 N VAL 265	13.670 0.073 13.955 1.00 22.25
	<b>ATOM</b>	825 CA VAL 265	13.754 -0.401 12.583 1.00 22.25
	ATOM	826 CB VAL 265	12.428 -1.038 12.086 1.00 25.31
	ATOM	827 CG1 VAL 265	12.079 -2.239 12.936 1.00 25.31
	ATOM	828 CG2 VAL 265	11.302 -0.030 12.091 1.00 25.31
25	ATOM	829 C VAL 265	14.208 0.707 11.639 1.00 22.25
	ATOM	830 O VAL 265	14.615 0.434 10.513 1.00 25.31
	ATOM	831 N ARG 266	14.124 1.955 12.092 1.00 26.45
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	ATOM	836 NE ARG 266	10.074 5.216 10.925 1.00 38.04
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	ATOM	838 NH1 ARG 266	10.237 6.038 13.075 1.00 38.04
35	<b>ATOM</b>	839 NH2 ARG 266	8.196 5.411 12.240 1.00 38.04
	ATOM	840 C ARG 266	15.957 3.531 11.729 1.00 26.45
	<b>ATOM</b>	841 O ARG 266	16.296 4.717 11.660 1.00 38.04
	ATOM	842 N TYR 267	16.733 2.590 12.251 1.00 24.87
	ATOM	843 CA TYR 267	18.083 2.888 12.700 1.00 24.87
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	<b>ATOM</b>	845 CG TYR 267	20.073 1.865 13.931 1.00 25.84
	ATOM	846 CD1 TYR 267	20.579 2.789 14.844 1.00 25.84
	ATOM	847 CE1 TYR 267	21.940 2.865 15.103 1.00 25.84
	ATOM	848 CD2 TYR 267	20.971 1.017 13.284 1.00 25.84
45	ATOM	849 CE2 TYR 267	22.331 1.085 13.536 1.00 25.84
	ATOM	850 CZ TYR 267	22.810 2.011 14.444 1.00 25.84
	ATOM	851 OH TYR 267	24.162 2.078 14.683 1.00 25.84
	ATOM	852 C TYR 267	18.999 3.009 11.488 1.00 24.87
	ATOM	853 O TYR 267	19.019 2.130 10.625 1.00 25.84
50	ATOM	854 N ASP 268	19.751 4.102 11.423 1.00 28.13

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     ATOM
            891 OG1 THR 273
                                20.723 8.808 12.491 1.00 30.96
     ATOM
                                21.967 9.924 14.228 1.00 30.96
            892 CG2 THR 273
     ATOM
                              19.701 6.829 14.442 1.00 27.60
            893 C THR 273
     ATOM
            894 O THR 273
                               19.633 5.883 13.650 1.00 30.96
40
     ATOM
            895 N LEU 274
                               18.696 7.192 15.232 1.00 20.89
     ATOM
            896 CA LEU 274
     ATOM
                               17.374 6.574 15.161 1.00 20.89
            897 CB LEU 274
                               16.862 6.193 16.555 1.00 22.48
     ATOM
            898 CG LEU 274
                               17.480 5.009 17.301 1.00 22.48
     ATOM
45
     ATOM
            899 CD1 LEU 274
                                16.798 4.866 18.650 1.00 22.48
                                17.317 3.736 16.497 1.00 22.48
            900 CD2 LEU 274
     ATOM
                               16.470 7.654 14.586 1.00 20.89
            901 C LEU 274
     ATOM
            902 O LEU 274
                               16.753 8.842 14.744 1.00 22.48
     ATOM
            903 N THR 275
                               15.393 7.258 13.922 1.00 27.89
     ATOM
     ATOM 904 CA THR 275
                               14.478 8.235 13.354 1.00 27.89
50
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	ATOM ATOM	905 CB THR 275 906 OG1 THR 275	14.325 8.045 11.832 1.00 37.64 15.622 7.983 11.228 1.00 37.64
	ATOM	907 CG2 THR 275	13.570 9.215 11.222 1.00 37.64
	ATOM	908 C THR 275	13.120 8.135 14.032 1.00 27.89
5	ATOM	909 O THR 275	12.493 7.081 14.019 1.00 37.64
	ATOM	910 N LEU 276	12.700 9.226 14.667 1.00 28.07
	ATOM	911 CA LEU 276	11.418 9.275 15.358 1.00 28.07
	ATOM	912 CB LEU 276	11.497 10.214 16.572 1.00 24.81
	ATOM	913 CG LEU 276	12.639 10.005 17.577 1.00 24.81
10	ATOM	914 CD1 LEU 276	12.450 10.929 18.769 1.00 24.81
	ATOM	915 CD2 LEU 276	12.692 8.558 18.038 1.00 24.81
	ATOM	916 C LEU 276	10.339 9.761 14.395 1.00 28.07
	ATOM	917 O LEU 276	10.533 10.760 13.691 1.00 24.81
	ATOM	918 N SER 277	9.232 9.027 14.331 1.00 29.24
15	ATOM	919 CA SER 277	8.106 9.357 13.458 1.00 29.24
	ATOM	920 CB SER 277	7.369 10.594 13.985 1.00 30.56
	ATOM	921 OG SER 277	6.845 10.358 15.283 1.00 30.56
	<b>ATOM</b>	922 C SER 277	8.533 9.569 12.005 1.00 29.24
	<b>ATOM</b>	923 O SER 277	7.902 10.326 11.263 1.00 30.56
20	<b>ATOM</b>	924 N GLY 278	9.619 8.908 11.618 1.00 34.41
	ATOM	925 CA GLY 278	10.135 9.024 10.263 1.00 34.41
	<b>ATOM</b>	926 C GLY 278	10.472 10.442 9.830 1.00 34.41
	ATOM	927 O GLY 278	10.516 10.725 8.631 1.00 44.04
	ATOM	928 N GLU 279	10.733 11.326 10.791 1.00 37.82
25	ATOM	929 CA GLU 279	11.056 12.717 10.479 1.00 37.82
	ATOM	930 CB GLU 279	9.808 13.600 10.612 1.00 70.24
	ATOM	931 CG GLU 279	9.202 13.631 12.014 1.00 70.24
	ATOM	932 CD GLU 279	8.028 14.593 12.141 1.00 70.24
	ATOM	933 OE1 GLU 279	8.028 15.406 13.093 1.00 70.24
30	ATOM	934 OE2 GLU 279	7.103 14.535 11.301 1.00 70.24
	ATOM	935 C GLU 279	12.192 13.321 11.300 1.00 37.82
	ATOM	936 O GLU 279	12.857 14.248 10.841 1.00 70.24
	ATOM	937 N MET 280	12.424 12.811 12.505 1.00 33.77
	ATOM	938 CA MET 280	13.482 13.360 13.344 1.00 33.77
35	ATOM	939 CB MET 280	12.903 13.848 14.674 1.00 33.89
	ATOM	940 CG MET 280	13.898 14.595 15.545 1.00 33.89
	ATOM	941 SD MET 280	13.350 14.740 17.256 1.00 33.89
	ATOM	942 CE MET 280	12.100 16.017 17.121 1.00 33.89
40	ATOM	943 C MET 280	14.620 12.383 13.613 1.00 33.77
40	ATOM	944 O MET 280	14.432 11.366 14.282 1.00 33.89
	ATOM	945 N ALA 281	15.797 12.690 13.080 1.00 30.24
	ATOM	946 CA ALA 281	16.972 11.852 13.287 1.00 30.24
	ATOM	947 CB ALA 281	17.937 11.998 12.120 1.00 25.10
15	ATOM	948 C ALA 281	17.631 12.309 14.587 1.00 30.24
45	ATOM	949 O ALA 281	18.008 13.477 14.718 1.00 25.10
	ATOM	950 N VAL 282	17.743 11.401 15.551 1.00 32.12
	ATOM	951 CA VAL 282	18.339 11.726 16.844 1.00 32.12 17.303 11.606 17.991 1.00 37.75
	ATOM ATOM	952 CB VAL 282	16.184 12.615 17.799 1.00 37.75
50	ATOM	953 CG1 VAL 282 954 CG2 VAL 282	16.184 12.615 17.799 1.00 37.75 16.739 10.193 18.055 1.00 37.75
50	ATOM	934 CG2 VAL 282	10./5 10.15 10.05 10.05 1.00 5/./5

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ATOM
            955 C VAL 282
                               19.543 10.852 17.181 1.00 32.12
     ATOM
            956 O VAL 282
                               19.614 9.690 16.778 1.00 37.75
            957 N LYS 283
                               20.491 11.428 17.913 1.00 26.82
     ATOM
            958 CA LYS 283
                               21.700 10.722 18.328 1.00 26.82
     ATOM
            959 CB LYS 283
                               22.894 11.679 18.342 1.00 57.25
     ATOM
            960 CG LYS 283
                               23.258 12.245 16.979 1.00 57.25
     ATOM
            961 CD LYS 283
                               24.282 13.361 17.105 1.00 57.25
     ATOM
            962 CE LYS 283
                               24.752 13.836 15.741 1.00 57.25
     ATOM
            963 NZ LYS 283
                               25.518 12.772 15.033 1.00 57.25
     ATOM
                              21.509 10.120 19.717 1.00 26.82
            964 C LYS 283
10
     ATOM
                               20,648 10,566 20,477 1.00 57.25
            965 O LYS 283
     ATOM
            966 N ARG 284
                               22.351 9.146 20.058 1.00 26.41
     ATOM
            967 CA ARG 284
                               22.297 8.457 21.351 1.00 26.41
     ATOM
     ATOM
            968 CB ARG 284
                                23.527 7.566 21.528 1.00 41.02
15
     ATOM
            969 CG ARG 284
                                23.715 6.539 20.440 1.00 41.02
            970 CD ARG 284
     ATOM
                                25.016 5.794 20.616 1.00 41.02
            971 NE ARG 284
                                25.145 4.730 19.630 1.00 41.02
     ATOM
            972 CZ ARG 284
                                24.759 3.475 19.831 1.00 41.02
     ATOM
            973 NH1 ARG 284
                                24.221 3.117 20.990 1.00 41.02
     ATOM
            974 NH2 ARG 284
                                24.886 2.584 18.859 1.00 41.02
20
     ATOM
            975 C ARG 284
                               22,200 9.399 22.543 1.00 26.41
     ATOM
            976 O ARG 284
                               21.296 9.278 23.370 1.00 41.02
     ATOM
            977 N GLU 285
     ATOM
                               23.152 10.321 22.634 1.00 33.23
            978 CA GLU 285
     ATOM
                               23.201 11.292 23.721 1.00 33.23
                               24.366 12.258 23.492 1.00 69.82
            979 CB GLU 285
25
     ATOM
                                24.485 13.359 24.533 1.00 69.82
            980 CG GLU 285
     ATOM
     ATOM
            981 CD GLU 285
                                25.079 14.636 23.964 1.00 69.82
     ATOM
            982 OE1 GLU 285
                                26.309 14.826 24.070 1.00 69.82
            983 OE2 GLU 285
     ATOM
                                24.309 15.453 23.409 1.00 69.82
30
     ATOM
            984 C GLU 285
                               21.898 12.082 23.823 1.00 33.23
            985 O GLU 285
     ATOM
                               21.336 12.239 24.907 1.00 69.82
            986 N GLN 286
     ATOM
                               21.414 12.551 22.677 1.00 28.07
     ATOM
            987 CA GLN 286
                               20.194 13.346 22.614 1.00 28.07
                                19.948 13.824 21.181 1.00 41.05
     ATOM
            988 CB GLN 286
35
     ATOM
            989 CG GLN 286
                                21.051 14.726 20.639 1.00 41.05
     ATOM
            990 CD GLN 286
                                20.808 15.154 19.202 1.00 41.05
           991 OEI GLN 286
                                20.783 14.322 18.293 1.00 41.05
     ATOM
     ATOM 992 NE2 GLN 286
                                20.635 16.452 18.990 1.00 41.05
     ATOM
           993 C GLN 286
                               18.955 12.642 23.162 1.00 28.07
            994 O GLN 286
                               18.281 13.174 24.048 1.00 41.05
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     ATOM
            995 N LEU 287
     ATOM
                               18.663 11.447 22.658 1.00 30.11
     ATOM
            996 CA LEU 287
                               17.492 10.705 23.116 1.00 30.11
            997 CB LEU 287
                               17.232 9.489 22.219 1.00 21.70
     ATOM
            998 CG LEU 287
                               15.859 8.821 22.357 1.00 21.70
     ATOM
            999 CD1 LEU 287
45
     ATOM
                                14.748 9.818 22.061 1.00 21.70
     ATOM 1000 CD2 LEU 287
                                15.763 7.628 21.421 1.00 21.70
     ATOM 1001 C LEU 287
                               17.641 10.277 24.577 1.00 30.11
     ATOM 1002 O LEU 287
                               16.655 10.212 25.320 1.00 21.70
     ATOM 1003 N LYS 288
                               18.878 10.015 24.992 1.00 20.72
     ATOM 1004 CA LYS 288
                               19.156 9.611 26.365 1.00 20.72
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20.626 9.213 26.514 1.00 43.14
    ATOM 1005 CB LYS 288
                               20.991 8.721 27.903 1.00 43.14
    ATOM 1006 CG LYS 288
                               22.374 8.102 27.931 1.00 43.14
    ATOM 1007 CD LYS 288
                               22.615 7.379 29.250 1.00 43.14
    ATOM 1008 CE LYS 288
                               23.866 6.568 29.224 1.00 43.14
    ATOM 1009 NZ LYS 288
                               18.819 10.742 27.331 1.00 20.72
    ATOM 1010 C LYS 288
                               18.027 10.566 28.261 1.00 43.14
    ATOM 1011 O LYS 288
                               19.380 11.917 27.067 1.00 33.64
    ATOM 1012 N ASN 289
    ATOM 1013 CA ASN 289
                                19.156 13.090 27.906 1.00 33.64
    ATOM 1014 CB ASN 289
                               20.190 14.173 27.590 1.00 35.61
10
    ATOM 1015 CG ASN 289
                                21.607 13.730 27.898 1.00 35.61
                                21.835 12.920 28.797 1.00 35.61
    ATOM 1016 OD1 ASN 289
    ATOM 1017 ND2 ASN 289
                                22.566 14.253 27.149 1.00 35.61
                               17.747 13.654 27.757 1.00 33.64
    ATOM 1018 C ASN 289
                               17.276 14.399 28.616 1.00 35.61
    ATOM 1019 O ASN 289
15
    ATOM 1020 N GLY 290
                               17.072 13.287 26.672 1.00 22.05
                               15.722 13.767 26.435 1.00 22.05
    ATOM 1021 CA GLY 290
                               14.688 13.247 27.416 1.00 22.05
    ATOM 1022 C GLY 290
    ATOM 1023 O GLY 290
                               13.550 13.710 27.420 1.00 29.95
                               15.072 12.276 28.239 1.00 24.91
    ATOM 1024 N GLY 291
20
    ATOM 1025 CA GLY 291
                               14.142 11.732 29.211 1.00 24.91
    ATOM 1026 C GLY 291
                               14.093 10.217 29.248 1.00 24.91
    ATOM 1027 O GLY 291
                               13.536 9.640 30.179 1.00 29.39
     ATOM 1028 N LEU 292
                               14.676 9.567 28.246 1.00 30.21
                               14.675 8.110 28.189 1.00 30.21
25
    ATOM 1029 CA LEU 292
    ATOM 1030 CB LEU 292
                                14.732 7.626 26.734 1.00 21.45
                                13.439 7.795 25.928 1.00 21.45
     ATOM 1031 CG LEU 292
     ATOM 1032 CD1 LEU 292
                                13.612 7.225 24.542 1.00 21.45
     ATOM 1033 CD2 LEU 292
                                12.296 7.087 26.630 1.00 21.45
                               15.785 7.461 29.013 1.00 30.21
     ATOM 1034 C LEU 292
30
                               15.645 6.324 29.473 1.00 21.45
     ATOM 1035 O LEU 292
                               16.885 8.180 29.205 1.00 16.29
     ATOM 1036 N GLY 293
     ATOM 1037 CA GLY 293
                                17.992 7.638 29.970 1.00 16.29
    ATOM 1038 C GLY 293
ATOM 1039 O GLY 293
                               18.534 6.374 29.332 1.00 16.29
                               18,763 6.334 28.122 1.00 25.88
35
                               18.689 5.322 30.130 1.00 33.05
     ATOM 1040 N VAL 294
                                19.211 4.050 29.635 1.00 33.05
     ATOM 1041 CA VAL 294
                                19.530 3.069 30.788 1.00 30.11
     ATOM 1042 CB VAL 294
                                20.718 3.577 31.582 1.00 30.11
     ATOM 1043 CG1 VAL 294
                               18.315 2.887 31.697 1.00 30.11
     ATOM 1044 CG2 VAL 294
40
                               18.302 3.361 28.617 1.00 33.05
     ATOM 1045 C VAL 294
                               18.768 2.545 27.817 1.00 30.11
     ATOM 1046 O VAL 294
     ATOM 1047 N VAL 295
                               17.014 3.699 28.635 1.00 18.14
                                16.056 3.118 27.698 1.00 18.14
     ATOM 1048 CA VAL 295
                                14.638 3.698 27.902 1.00 28.34
45
     ATOM 1049 CB VAL 295
                                13.668 3.099 26.893 1.00 28.34
     ATOM 1050 CG1 VAL 295
                                14.159 3.431 29.317 1.00 28.34
     ATOM 1051 CG2 VAL 295
                               16.521 3.415 26.275 1.00 18.14
     ATOM 1052 C VAL 295
                               16.395 2.577 25.383 1.00 28.34
     ATOM 1053 O VAL 295
                               17.091 4.601 26.085 1.00 20.84
     ATOM 1054 N SER 296
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ATOM 1055 CA SER 296
                               17.596 5.028 24.785 1.00 20.84
                               18.160 6.446 24.884 1.00 25.61
    ATOM 1056 CB SER 296
                                18.615 6.911 23.627 1.00 25.61
    ATOM 1057 OG SER 296
    ATOM 1058 C SER 296
                               18.687 4.074 24.307 1.00 20.84
    ATOM 1059 O SER 296
                               18.723 3.691 23.133 1.00 25.61
                               19.571 3.691 25.224 1.00 28.08
    ATOM 1060 N ASP 297
                               20.660 2.777 24.904 1.00 28.08
    ATOM 1061 CA ASP 297
                               21.555 2.552 26.129 1.00 51.15
    ATOM 1062 CB ASP 297
                               22.207 3.835 26.629 1.00 51.15
    ATOM 1063 CG ASP 297
                                22.508 4.725 25.804 1.00 51.15
    ATOM 1064 OD1 ASP 297
10
    ATOM 1065 OD2 ASP 297
                                22.425 3.948 27.855 1.00 51.15
                               20.079 1.450 24.434 1.00 28.08
    ATOM 1066 C ASP 297
    ATOM 1067 O ASP 297
                               20.549 0.869 23.456 1.00 51.15
    ATOM 1068 N ALA 298
                               19.024 1.006 25.111 1.00 26.12
    ATOM 1069 CA ALA 298
                                18.357 -0.245 24.778 1.00 26.12
15
                                17.253 -0.530 25.787 1.00 18.80
    ATOM 1070 CB ALA 298
    ATOM 1071 C ALA 298
                               17.790 -0.223 23.356 1.00 26.12
                               18.014 -1.154 22.575 1.00 18.80
    ATOM 1072 O ALA 298
    ATOM 1073 N ILE 299
                              17.078 0.848 23.013 1.00 17.42
    ATOM 1074 CA ILE 299
                               16.483 0.979 21.686 1.00 17.42
20
    ATOM 1075 CB ILE 299
                               15.559 2.211 21.597 1.00 16.69
    ATOM 1076 CG2 ILE 299
                               14.845 2.238 20.253 1.00 16.69
    ATOM 1077 CG1 ILE 299
                               14.515 2.149 22.712 1.00 16.69
                               13.713 3.406 22.872 1.00 16.69
    ATOM 1078 CD1 ILE 299
                              17.563 1.042 20.609 1.00 17.42
    ATOM 1079 C ILE 299
25
                              17.416 0.443 19.542 1.00 16.69
    ATOM 1080 O ILE 299
    ATOM 1081 N PHE 300
                               18.652 1.752 20.889 1.00 14.46
    ATOM 1082 CA PHE 300
                               19.751 1.851 19.935 1.00 14.46
    ATOM 1083 CB PHE 300
                                20.804 2.854 20.409 1.00 24.01
    ATOM 1084 CG PHE 300
                                20.656 4.221 19.801 1.00 24.01
30
    ATOM 1085 CD1 PHE 300
                                19.904 5.204 20.435 1.00 24.01
    ATOM 1086 CD2 PHE 300
                                21.271 4.526 18.591 1.00 24.01
    ATOM 1087 CE1 PHE 300
                                19.766 6.472 19.873 1.00 24.01
    ATOM 1088 CE2 PHE 300
                                21.140 5.791 18.020 1.00 24.01
    ATOM 1089 CZ PHE 300
                               20.385 6.765 18.663 1.00 24.01
35
    ATOM 1090 C PHE 300
                               20.383 0.480 19.726 1.00 14.46
    ATOM 1091 O PHE 300
                               20.696 0.102 18.596 1.00 24.01
    ATOM 1092 N GLU 301
                               20.547 -0.270 20.813 1.00 21.61
    ATOM 1093 CA GLU 301
                               21.123 -1.609 20.744 1.00 21.61
    ATOM 1094 CB GLU 301
                                21.289 -2.192 22.143 1.00 23.89
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    ATOM 1095 C GLU 301
                               20.211 -2.498 19.904 1.00 21.61
     ATOM 1096 O GLU 301
                               20.681 -3.251 19.043 1.00 23.89
                               18.906 -2.390 20.140 1.00 14.43
    ATOM 1097 N LEU 302
                               17.922 -3.168 19.399 1.00 14.43
    ATOM 1098 CA LEU 302
                                16.512 -2.872 19.912 1.00 23.43
    ATOM 1099 CB LEU 302
45
                                15.350 -3.669 19.312 1.00 23.43
    ATOM 1100 CG LEU 302
                               15.459 -5.140 19.688 1.00 23.43
    ATOM 1101 CD1 LEU 302
     ATOM 1102 CD2 LEU 302
                               14.035 -3.094 19.804 1.00 23.43
    ATOM 1103 C LEU 302
                               18.027 -2.812 17.917 1.00 14.43
                               18.089 -3.697 17.066 1.00 23.43
    ATOM 1104 O LEU 302
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	ATOM	1105 N GLY 303	18.098 -1.515 17.625 1.00 15.17
	ATOM	1106 CA GLY 303	18.208 -1.056 16.251 1.00 15.17
	ATOM	1107 C GLY 303	19.411 -1.640 15.530 1.00 15.17
	ATOM	1108 O GLY 303	19.290 -2.137 14.406 1.00 27.67
5	ATOM	1109 N LYS 304	20.570 -1.594 16.182 1.00 19.04
	ATOM	1110 CA LYS 304	21.802 -2.127 15.605 1.00 19.04
	ATOM	1111 CB LYS 304	22.979 -1.975 16.577 1.00 56.94
	ATOM	1112 CG LYS 304	23.496 -0.556 16.741 1.00 56.94
	ATOM	1113 CD LYS 304	24.811 -0.524 17.516 1.00 56.94
10	ATOM	1114 CE LYS 304	24.634 -0.965 18.968 1.00 56.94
10	ATOM	1115 NZ LYS 304	23.838 0.008 19.778 1.00 56.94
	ATOM	1116 C LYS 304	21.653 -3.596 15.229 1.00 19.04
	ATOM	1117 O LYS 304	21.974 -3.993 14.107 1.00 56.94
	ATOM	1117 O E13 304 1118 N SER 305	21.146 -4.394 16.164 1.00 24.46
15	ATOM	1119 CA SER 305	20.965 -5.822 15.932 1.00 24.46
15	ATOM	1119 CA SER 305	20.610 -6.533 17.240 1.00 37.46
	ATOM	1121 OG SER 305	19.444 -5.984 17.827 1.00 37.46
		1121 OG SER 305	19.926 -6.128 14.853 1.00 24.46
	ATOM		20.146 -6.996 14.006 1.00 37.46
20	ATOM		18.819 -5.390 14.858 1.00 25.47
20	ATOM		17.753 -5.592 13.881 1.00 25.47
	ATOM	1125 CA LEU 306	16.525 -4.746 14.224 1.00 15.99
	ATOM	1126 CB LEU 306	
	ATOM	1127 CG LEU 306	15.700 -5.190 15.432 1.00 15.99
0.5	ATOM	1128 CD1 LEU 306	14.504 -4.271 15.600 1.00 15.99
25	ATOM	1129 CD2 LEU 306	15.244 -6.624 15.247 1.00 15.99
	ATOM	1130 C LEU 306	18.174 -5.330 12.439 1.00 25.47
	ATOM	1131 O LEU 306	17.596 -5.902 11.513 1.00 15.99
	ATOM	1132 N SER 307	19.182 -4.482 12.247 1.00 24.28
	ATOM	1133 CA SER 307	19.670 -4.160 10.907 1.00 24.28
30	ATOM	1134 CB SER 307	20.910 -3.263 10.989 1.00 40.92
	ATOM	1135 OG SER 307	20.617 -2.028 11.622 1.00 40.92
	ATOM	1136 C SER 307	19.995 -5.422 10.107 1.00 24.28
	ATOM	1137 O SER 307	19.625 -5.535 8.936 1.00 40.92
	ATOM	1138 N ALA 308	20.644 -6.383 10.761 1.00 30.97
35	ATOM	1139 CA ALA 308	21.027 -7.640 10.124 1.00 30.97
	ATOM	1140 CB ALA 308	22.004 -8.399 11.013 1.00 37.84
	ATOM	1141 C ALA 308	19.830 -8.528 9.779 1.00 30.97
	ATOM	1142 O ALA 308	19.897 -9.336 8.853 1.00 37.84
	ATOM	1143 N PHE 309	18.737 -8.372 10.520 1.00 22.78
40	ATOM	1144 CA PHE 309	17.533 -9.166 10.292 1.00 22.78
	ATOM	1145 CB PHE 309	16.571 -9.037 11.477 1.00 30.14
	ATOM	1146 CG PHE 309	17.032 -9.751 12.716 1.00 30.14
	<b>ATOM</b>	1147 CD1 PHE 309	16.299 -10.809 13.236 1.00 30.14
	ATOM	1148 CD2 PHE 309	18.204 -9.372 13.359 1.00 30.14
45	<b>ATOM</b>	1149 CE1 PHE 309	16.725 -11.481 14.378 1.00 30.14
	ATOM	1150 CE2 PHE 309	18.640 -10.038 14.503 1.00 30.14
	ATOM	1151 CZ PHE 309	17.896 -11.094 15.013 1.00 30.14
	ATOM	1152 C PHE 309	16.818 -8.813 8.990 1.00 22.78
	ATOM	1153 O PHE 309	16.068 -9.631 8.451 1.00 30.14
50	ATOM	1154 N ASN 310	17.051 -7.598 8.496 1.00 35.30

PCT/US98/25296

# WO 99/26966

	ATOM	1155 CA ASN 310	16.441 -7.109 7.255 1.00 35.30
	ATOM	1156 CB ASN 310	17.109 -7.760 6.037 1.00 28.28
	ATOM	1157 C ASN 310	14.929 -7.339 7.229 1.00 35.30
	ATOM	1158 O ASN 310	14.395 -7.970 6.312 1.00 28.28
5	ATOM	1159 N LEU 311	14.249 -6.831 8.251 1.00 27.52
	ATOM	1160 CA LEU 311	12.803 -6.979 8.369 1.00 27.52
	ATOM	1161 CB LEU 311	12.351 -6.630 9.788 1.00 22.62
	ATOM	1162 CG LEU 311	12.950 -7.396 10.968 1.00 22.62
	ATOM	1163 CD1 LEU 311	12.360 -6.864 12.268 1.00 22.62
10	ATOM	1164 CD2 LEU 311	12.672 -8.881 10.821 1.00 22.62
	ATOM	1165 C LEU 311	12.060 -6.085 7.382 1.00 27.52
	ATOM	1166 O LEU 311	12.519 -4.986 7.067 1.00 22.62
	ATOM	1167 N ASP 312	10.918 -6.563 6.892 1.00 16.74
	ATOM	1168 CA ASP 312	10.095 -5.789 5.968 1.00 16.74
15	ATOM	1169 CB ASP 312	9.803 -6.578 4.673 1.00 16.35
	ATOM	1170 CG ASP 312	8.924 -7.814 4.888 1.00 16.35
	ATOM	1171 OD1 ASP 312	8.591 -8.168 6.037 1.00 16.35
	ATOM	1172 OD2 ASP 312	8.559 -8.446 3.876 1.00 16.35
	ATOM	1173 C ASP 312	8.808 -5.354 6.678 1.00 16.74
20	ATOM	1174 O ASP 312	8.535 -5.798 7.797 1.00 16.35
	ATOM	1175 N ASP 313	8.007 -4.520 6.019 1.00 5.43
	ATOM		6.758 -4.016 6.592 1.00 5.43
	ATOM	1177 CB ASP 313	5.974 -3.201 5.559 1.00 31.80
	ATOM	1178 CG ASP 313	6.670 -1.906 5.183 1.00 31.80
25	ATOM	1179 OD1 ASP 313	7.392 -1.340 6.033 1.00 31.80
	ATOM	1180 OD2 ASP 313	6.493 -1.452 4.032 1.00 31.80
	ATOM	1181 C ASP 313	5.849 -5.081 7.189 1.00 5.43
	ATOM	1182 O ASP 313	5.216 -4.849 8.221 1.00 31.80
	ATOM	1183 N THR 314	5.777 -6.238 6.543 1.00 12.98
30	ATOM	1184 CA THR 314	4.934 -7.327 7.022 1.00 12.98
-	ATOM	1185 CB THR 314	4.825 -8.441 5.968 1.00 18.90
	ATOM	1186 OG1 THR 314	4.249 -7.904 4.769 1.00 18.90
	ATOM	1187 CG2 THR 314	3.960 -9.578 6.477 1.00 18.90
	ATOM	1188 C THR 314	5.426 -7.910 8.349 1.00 12.98
35	ATOM	1189 O THR 314	4.636 -8.124 9.268 1.00 18.90
55	ATOM	1190 N GLU 315	6.731 -8.135 8.457 1.00 9.13
	ATOM	1191 CA GLU 315	7.316 -8.685 9.675 1.00 9.13
	ATOM	1192 CB GLU 315	8.771 -9.078 9.427 1.00 11.49
	ATOM	1193 CG GLU 315	8.870 -10.323 8.562 1.00 11.49
40	ATOM	1194 CD GLU 315	10.233 -10.544 7.945 1.00 11.49
	ATOM	1195 OE1 GLU 315	10.964 -9.561 7.705 1.00 11.49
	ATOM	1196 OE2 GLU 315	10.558 -11.715 7.669 1.00 11.49
	ATOM	1197 C GLU 315	7.180 -7.720 10.847 1.00 9.13
	ATOM	1198 O GLU 315	6.863 -8.131 11.967 1.00 11.49
45	ATOM	1199 N VAL 316	7.376 -6.433 10.575 1.00 9.46
73	ATOM	1200 CA VAL 316	7.240 -5.406 11.602 1.00 9.46
	ATOM	1200 CA VAL 316	7.655 -4.015 11.063 1.00 7.95
	ATOM	1201 CB VAL 316	7.434 -2.941 12.124 1.00 7.95
	ATOM	1202 CG1 VAL 316	9.112 -4.037 10.625 1.00 7.95
50	ATOM	1204 C VAL 316	5.777 -5.365 12.051 1.00 9.46
50	VIONI	1204 C AUD 210	J.,,, -J.JOJ 12.031 1.00 7.70

WO 99/26966

PCT/US98/25296

	ATOM	1205 O VAL 316	5.484 -5.300 13.247 1.00 7.95
	ATOM	1206 N ALA 317	4.866 -5.438 11.083 1.00 5.52
	ATOM	1207 CA ALA 317	3.434 -5.417 11.355 1.00 5.52
	ATOM	1208 CB ALA 317	2.656 -5.415 10.054 1.00 10.98
5	ATOM	1209 C ALA 317	3.002 -6.595 12.225 1.00 5.52
	ATOM	1210 O ALA 317	2.317 -6.412 13.230 1.00 10.98
	ATOM	1211 N LEU 318	3.411 -7.799 11.838 1.00 8.62
	ATOM	1212 CA LEU 318	3.067 -9.003 12.584 1.00 8.62
	ATOM	1213 CB LEU 318	3.523 -10.249 11.825 1.00 10.49
10	ATOM	1214 CG LEU 318	2.770 -10.494 10.514 1.00 10.49
	ATOM	1215 CD1 LEU 318	3.376 -11.664 9.769 1.00 10.49
	ATOM	1216 CD2 LEU 318	1.297 -10.741 10.799 1.00 10.49
	ATOM	1217 C LEU 318	3.674 -8.971 13.978 1.00 8.62
	ATOM	1218 O LEU 318	3.047 -9.407 14.945 1.00 10.49
15	ATOM	1219 N LEU 319	4.885 -8.435 14.082 1.00 9.43
	ATOM	1220 CA LEU 319	5.560 -8.325 15.366 1.00 9.43
	ATOM	1221 CB LEU 319	6.975 -7.773 15.173 1.00 24.05
	ATOM	1222 CG LEU 319	7.901 -7.680 16.389 1.00 24.05
	ATOM	1223 CD1 LEU 319	7.889 -8.977 17.182 1.00 24.05
20	ATOM	1224 CD2 LEU 319	9.310 -7.356 15.922 1.00 24.05
	ATOM	1225 C LEU 319	4.731 -7.404 16.259 1.00 9.43
	ATOM	1226 O LEU 319	4.456 -7.731 17.416 1.00 24.05
	ATOM	1227 N GLN 320	4.287 -6.282 15.699 1.00 8.67
	<b>ATOM</b>	1228 CA GLN 320	3.467 -5.325 16.437 1.00 8.67
25	ATOM	1229 CB GLN 320	3.151 -4.102 15.573 1.00 10.94
	ATOM	1230 CG GLN 320	4.361 -3.256 15.218 1.00 10.94
	ATOM	1231 CD GLN 320	4.025 -2.045 14.359 1.00 10.94
	ATOM	1232 OE1 GLN 320	4.889 -1.217 14.082 1.00 10.94
	ATOM	1233 NE2 GLN 320	2.773 -1.940 13.924 1.00 10.94
30	ATOM	1234 C GLN 320	2.169 -5.984 16.895 1.00 8.67
	ATOM	1235 O GLN 320	1.708 -5.751 18.013 1.00 10.94
	ATOM	1236 N ALA 321	1.586 -6.806 16.028 1.00 9.21
	ATOM	1237 CA ALA 321	0.349 -7.513 16.342 1.00 9.21
	ATOM	1238 CB ALA 321	-0.136 -8.283 15.129 1.00 12.83
35	ATOM	1239 C ALA 321	0.558 -8.460 17.523 1.00 9.21
	ATOM	1240 O ALA 321	-0.315 -8.591 18.382 1.00 12.83
	ATOM	1241 N VAL 322	1.718 -9.111 17.566 1.00 9.10
	ATOM	1242 CA VAL 322	2.043 -10.030 18.651 1.00 9.10
	ATOM	1243 CB VAL 322	3.340 -10.827 18.352 1.00 15.92
40	ATOM	1244 CG1 VAL 322	3.783 -11.614 19.575 1.00 15.92
	ATOM	1245 CG2 VAL 322	3.106 -11.780 17.194 1.00 15.92
	ATOM	1246 C VAL 322	2.192 -9.256 19.960 1.00 9.10
	ATOM	1247 O VAL 322	1.707 -9.691 21.003 1.00 15.92
	ATOM	1248 N LEU 323	2.856 -8.106 19.893 1.00 11.07
45	ATOM	1249 CA LEU 323	3.062 -7.257 21.064 1.00 11.07
	ATOM	1250 CB LEU 323	3.959 -6.070 20.705 1.00 16.31
	ATOM	1251 CG LEU 323	5,377 -6.393 20.229 1.00 16.31
	ATOM	1252 CD1 LEU 323	6.039 -5.149 19.669 1.00 16.31
	ATOM	1253 CD2 LEU 323	6.187 -6.966 21.375 1.00 16.31
50	ATOM	1254 C LEU 323	1.729 -6.742 21.595 1.00 11.07
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ATOM 1255 O LEU 323
                                1.523 -6.650 22.803 1.00 16.31
    ATOM 1256 N LEU 324
                                0.827 -6.413 20.677 1.00 13.48
     ATOM 1257 CA LEU 324
                                -0.494 -5.900 21.015 1.00 13.48
    ATOM 1258 CB LEU 324
                                -1.185 -5.383 19.752 1.00 15.92
                                -2.607 -4.837 19.889 1.00 15.92
    ATOM 1259 CG LEU 324
                                 -2.602 -3.547 20.692 1.00 15.92
    ATOM 1260 CD1 LEU 324
                                 -3.182 -4.598 18.511 1.00 15.92
    ATOM 1261 CD2 LEU 324
                                -1.393 -6.924 21.707 1.00 13.48
    ATOM 1262 C LEU 324
    ATOM 1263 O LEU 324
                                -1.896 -6.678 22.802 1.00 15.92
    ATOM 1264 N MET 325
                                -1.593 -8.074 21.072 1.00 11.47
10
                                 -2.458 -9.111 21.631 1.00 11.47
    ATOM 1265 CA MET 325
    ATOM 1266 CB MET 325
                                 -2.959 -10.043 20.520 1.00 22.90
                                 -3.689 -9.347 19.375 1.00 22.90
    ATOM 1267 CG MET 325
    ATOM 1268 SD MET 325
                                -5.052 -8.287 19.908 1.00 22.90
    ATOM 1269 CE MET 325
                                -6.284 -9.475 20.353 1.00 22.90
15
    ATOM 1270 C MET 325
ATOM 1271 O MET 325
                                -1.814 -9.932 22.752 1.00 11.47
                                -1.899 -11.160 22.758 1.00 22.90
    ATOM 1272 N SER 326
                               -1.193 -9.256 23.711 1.00 30.07
    ATOM 1273 CA SER 326
                                -0.543 -9.936 24.826 1.00 30.07
                                 0.723 -9.175 25.239 1.00 32.79
20
    ATOM 1274 CB SER 326
                                1.283 -9.699 26.433 1.00 32.79
    ATOM 1275 OG SER 326
    ATOM 1276 C SER 326
                               -1.492 -10.061 26.014 1.00 30.07
    ATOM 1277 O SER 326
                               -2.343 -9.198 26.235 1.00 32.79
                                -1.347 -11.143 26.773 1.00 29.08
    ATOM 1278 N THR 327
                                -2.179 -11.368 27.948 1.00 29.08
25
    ATOM 1279 CA THR 327
                                -2.705 -12.817 27.998 1.00 36.96
    ATOM 1280 CB THR 327
    ATOM 1281 OG1 THR 327
                                 -1.612 -13.734 27.856 1.00 36.96
    ATOM 1282 CG2 THR 327
                                 -3.716 -13.055 26.890 1.00 36.96
    ATOM 1283 C THR 327
                                -1.426 -11.049 29.239 1.00 29.08
    ATOM 1284 O THR 327
                                -1.930 -11.295 30.333 1.00 36.96
30
                                -0.214 -10.513 29.111 1.00 38.93
    ATOM 1285 N ASP 328
    ATOM 1286 CA ASP 328
                                 0.596 -10.152 30.273 1.00 38.93
    ATOM 1287 CB ASP 328
                                 2.082 -10.089 29.899 1.00 85.70
    ATOM 1288 CG ASP 328
                                 2.660 -11.451 29.556 1.00 85.70
                                 3.388 -11.554 28.542 1.00 85.70
35
    ATOM 1289 OD1 ASP 328
    ATOM 1290 OD2 ASP 328
                                 2.393 -12.418 30.303 1.00 85.70
    ATOM 1291 C ASP 328
                                0.148 -8.810 30.845 1.00 38.93
    ATOM 1292 O ASP 328
                                0.962 -7.911 31.061 1.00 85.70
    ATOM 1293 N ARG 329
                                -1.154 -8.673 31.070 1.00 28.95
                                 -1.716 -7.445 31.608 1.00 28.95
    ATOM 1294 CA ARG 329
40
    ATOM 1295 CB ARG 329
                                 -2.390 -6.612 30.509 1.00 38.88
    ATOM 1296 CG ARG 329
                                 -1.449 -5.887 29.554 1.00 38.88
    ATOM 1297 CD ARG 329
                                 -1.107 -6.739 28.347 1.00 38.88
                                 -0.322 -6.005 27.356 1.00 38.88
    ATOM 1298 NE ARG 329
    ATOM 1299 CZ ARG 329
                                 1.006 -5.936 27.351 1.00 38.88
45
    ATOM 1300 NH1 ARG 329
                                  1.713 -6.552 28.290 1.00 38.88
    ATOM 1301 NH2 ARG 329
                                  1.631 -5.270 26.391 1.00 38.88
    ATOM 1302 C ARG 329
                                -2.745 -7.790 32.672 1.00 28.95
    ATOM 1303 O ARG 329
                                -3.279 -8.898 32.696 1.00 38.88
     ATOM 1304 N SER 330
                               -3.029 -6.829 33.542 1.00 42.07
50
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ATOM 1305 CA SER 330
                                -3.999 -7.025 34.607 1.00 42.07
     ATOM 1306 CB SER 330
                                -3.488 -6.399 35.899 1.00 37.35
     ATOM 1307 C SER 330
                               -5.340 -6.413 34.220 1.00 42.07
     ATOM 1308 O SER 330
                               -5.386 -5.382 33.550 1.00 37.35
     ATOM 1309 N GLY 331
                                -6.424 -7.085 34.598 1.00 26.57
     ATOM 1310 CA GLY 331
                                -7.754 -6.572 34.318 1.00 26.57
                                -8.404 -6.915 32.991 1.00 26.57
     ATOM 1311 C GLY 331
     ATOM 1312 O GLY 331
                                -9.462 -6.371 32.671 1.00 30.06
     ATOM 1313 N LEU 332
                               -7.797 -7.807 32.214 1.00 31.47
     ATOM 1314 CA LEU 332
                                -8.374 -8.189 30.928 1.00 31.47
10
     ATOM 1315 CB LEU 332
                                -7.351 -8.933 30.065 1.00 23.83
    ATOM 1316 CG LEU 332
                                -6.261 -8.076 29.425 1.00 23.83
     ATOM 1317 CD1 LEU 332
                                 -5.296 -8.960 28.652 1.00 23.83
     ATOM 1318 CD2 LEU 332
                                 -6.897 -7.041 28.509 1.00 23.83
15
     ATOM 1319 C LEU 332
                               -9.630 -9.039 31.091 1.00 31.47
     ATOM 1320 O LEU 332
                               -9.665 -9.969 31.895 1.00 23.83
     ATOM 1321 N LEU 333
                               -10.659 -8.702 30.321 1.00 27.66
                                -11.927 -9.422 30.351 1.00 27.66
     ATOM 1322 CA LEU 333
                                -13.072 -8.500 29.918 1.00 49.79
     ATOM 1323 CB LEU 333
                                -13.416 -7.312 30.820 1.00 49.79
20
     ATOM 1324 CG LEU 333
     ATOM 1325 CD1 LEU 333
                                -14.328 -6.339 30.083 1.00 49.79
     ATOM 1326 CD2 LEU 333
                                -14.072 -7.803 32.104 1.00 49.79
     ATOM 1327 C LEU 333
                               -11.904 -10.663 29.456 1.00 27.66
     ATOM 1328 O LEU 333
                               -12.117 -11.780 29.919 1.00 49.79
     ATOM 1329 N CYS 334
                               -11.616 -10.464 28.174 1.00 29.56
25
                                -11.583 -11.566 27.220 1.00 29.56
    ATOM 1330 CA CYS 334
                                -12.134 -11.106 25.865 1.00 47.01
     ATOM 1331 CB CYS 334
    ATOM 1332 SG CYS 334
                               -13.888 -10.657 25.883 1.00 47.01
                               -10.187 -12.161 27.050 1.00 29.56
     ATOM 1333 C CYS 334
     ATOM 1334 O CYS 334
                               -9.652 -12.202 25.942 1.00 47.01
30
     ATOM 1335 N VAL 335
                                -9.617 -12.655 28.147 1.00 30.69
    ATOM 1336 CA VAL 335
                                -8.280 -13.250 28.132 1.00 30.69
    ATOM 1337 CB VAL 335
                                -7.913 -13.844 29.514 1.00 32.18
    ATOM 1338 CG1 VAL 335
                                 -6.517 -14.456 29.480 1.00 32.18
35
    ATOM 1339 CG2 VAL 335
                                -7.988 -12.768 30.584 1.00 32.18
     ATOM 1340 C VAL 335
                               -8.120 -14.340 27.068 1.00 30.69
     ATOM 1341 O VAL 335
                                -7.149 -14.337 26.309 1.00 32.18
     ATOM 1342 N ASP 336
                               -9.079 -15.260 27.012 1.00 30.13
    ATOM 1343 CA ASP 336
                                -9.040 -16.360 26.052 1.00 30.13
    ATOM 1344 CB ASP 336
                               -10.218 -17.311 26.284 1.00 63.22
40
    ATOM 1345 CG ASP 336
                                -10.178 -18.528 25.370 1.00 63.22
    ATOM 1346 OD1 ASP 336
                                -11.119 -18.700 24.565 1.00 63.22
    ATOM 1347 OD2 ASP 336
                                 -9.205 -19.311 25.452 1.00 63.22
                               -9.012 -15.903 24.594 1.00 30.13
     ATOM 1348 C ASP 336
                               -8.156 -16.339 23.823 1.00 63.22
45
    ATOM 1349 O ASP 336
                               -9.944 -15.027 24.223 1.00 26.63
    ATOM 1350 N LYS 337
    ATOM 1351 CA LYS 337
                               -10.024 -14.515 22.856 1.00 26.63
    ATOM 1352 CB LYS 337
                               -11.172 -13.516 22.729 1.00 21.38
    ATOM 1353 C LYS 337
                               -8.706 -13.865 22.438 1.00 26.63
    ATOM 1354 O LYS 337
                               -8.204 -14.110 21.338 1.00 21.38
50
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ATOM 1355 N ILE 338
                               -8.141 -13.060 23.334 1.00 24.65
                               -6.879 -12.376 23.078 1.00 24.65
    ATOM 1356 CA ILE 338
                               -6.543 -11.380 24.215 1.00 20.45
    ATOM 1357 CB ILE 338
                                -5.198 -10.719 23.966 1.00 20.45
    ATOM 1358 CG2 ILE 338
                                -7.632 -10.308 24.308 1.00 20.45
    ATOM 1359 CG1 ILE 338
                                -7.479 -9.374 25.486 1.00 20.45
    ATOM 1360 CD1 ILE 338
                               -5.744 -13.388 22.911 1.00 24.65
    ATOM 1361 C ILE 338
                               -4.948 -13.288 21.974 1.00 20.45
    ATOM 1362 O ILE 338
                                -5.700 -14.383 23.795 1.00 35.34
    ATOM 1363 N GLU 339
                                -4.673 -15.422 23.745 1.00 35.34
    ATOM 1364 CA GLU 339
10
                                -4.836 -16.388 24.916 1.00 29.51
    ATOM 1365 CB GLU 339
    ATOM 1366 C GLU 339
                                -4.744 -16.180 22.421 1.00 35.34
                                -3.720 -16.421 21.777 1.00 29.51
    ATOM 1367 O GLU 339
                               -5.959 -16.536 22.009 1.00 24.19
    ATOM 1368 N LYS 340
    ATOM 1369 CA LYS 340
                                -6.168 -17.256 20.755 1.00 24.19
15
    ATOM 1370 CB LYS 340
                                -7.627 -17.671 20.624 1.00 23.97
                               -5.754 -16.377 19.576 1.00 24.19
    ATOM 1371 C LYS 340
                               -5.197 -16.860 18.586 1.00 23.97
    ATOM 1372 O LYS 340
                               -6.000 -15.079 19.708 1.00 16.85
    ATOM 1373 N SER 341
    ATOM 1374 CA SER 341
                                -5.651 -14.115 18.676 1.00 16.85
20
                                -6.223 -12.744 19.033 1.00 26.59
    ATOM 1375 CB SER 341
                                -5.852 -11.765 18.080 1.00 26.59
    ATOM 1376 OG SER 341
                               -4.137 -14.026 18.500 1.00 16.85
     ATOM 1377 C SER 341
     ATOM 1378 O SER 341
                               -3.638 -14.042 17.374 1.00 26.59
                                -3.406 -13.932 19.608 1.00 17.35
     ATOM 1379 N GLN 342
25
                                -1.952 -13.845 19.537 1.00 17.35
    ATOM 1380 CA GLN 342
                                -1.337 -13.597 20.913 1.00 30.07
    ATOM 1381 CB GLN 342
     ATOM 1382 CG GLN 342
                                 0.140 -13.245 20.832 1.00 30.07
     ATOM 1383 CD GLN 342
                                 0.811 -13.196 22.182 1.00 30.07
                                 0.884 -14.201 22.884 1.00 30.07
30
     ATOM 1384 OE1 GLN 342
     ATOM 1385 NE2 GLN 342
                                  1.318 -12.030 22.548 1.00 30.07
                                -1.368 -15.118 18.944 1.00 17.35
     ATOM 1386 C GLN 342
     ATOM 1387 O GLN 342
                                -0.405 -15.066 18.178 1.00 30.07
     ATOM 1388 N GLU 343
                                -1.949 -16.260 19.303 1.00 18.35
35
     ATOM 1389 CA GLU 343
                                -1.489 -17.546 18.791 1.00 18.35
     ATOM 1390 CB GLU 343
                                 -2.308 -18.676 19.394 1.00 16.98
                                -1.603 -17.560 17.267 1.00 18.35
     ATOM 1391 C GLU 343
                                -0.699 -18.026 16.568 1.00 16.98
     ATOM 1392 O GLU 343
                                -2.706 -17.017 16.761 1.00 14.83
     ATOM 1393 N ALA 344
                                 -2.946 -16.948 15.324 1.00 14.83
     ATOM 1394 CA ALA 344
                                 -4.327 -16.376 15.049 1.00 19.42
     ATOM 1395 CB ALA 344
     ATOM 1396 C ALA 344
                                -1.872 -16.102 14.640 1.00 14.83
                                -1.311 -16.507 13.619 1.00 19.42
     ATOM 1397 O ALA 344
     ATOM 1398 N TYR 345
                                -1.586 -14.934 15.211 1.00 13.10
                                 -0.569 -14.041 14.665 1.00 13.10
     ATOM 1399 CA TYR 345
45
                                 -0.573 -12.697 15.393 1.00 2.00
     ATOM 1400 CB TYR 345
                                 -1.670 -11.767 14.938 1.00 2.00
     ATOM 1401 CG TYR 345
     ATOM 1402 CD1 TYR 345
                                 -2.707 -11.409 15.794 1.00 2.00
                                 -3.722 -10.562 15.377 1.00 2.00
     ATOM 1403 CE1 TYR 345
                                 -1.674 -11.248 13.647 1.00 2.00
     ATOM 1404 CD2 TYR 345
50
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	ATOM	1405 CE2 TYR 345	-2.683 -10.398 13.219 1.00 2.00
	ATOM	1406 CZ TYR 345	-3.706 -10.061 14.087 1.00 2.00
	ATOM	1407 OH TYR 345	-4.722 -9.233 13.669 1.00 2.00
	ATOM	1408 C TYR 345	0.818 -14.666 14.732 1.00 13.10
5	ATOM	1409 O TYR 345	1.614 -14.504 13.811 1.00 2.00
	ATOM	1410 N LEU 346	1.101 -15.387 15.813 1.00 12.59
	ATOM	1411 CA LEU 346	2.396 -16.041 15.976 1.00 12.59
	ATOM	1412 CB LEU 346	2.498 -16.715 17.347 1.00 22.61
	ATOM	1413 CG LEU 346	2.899 -15.799 18.504 1.00 22.61
10	ATOM	1414 CD1 LEU 346	2.717 -16.511 19.830 1.00 22.61
	<b>ATOM</b>	1415 CD2 LEU 346	4.341 -15.357 18.324 1.00 22.61
	ATOM	1416 C LEU 346	2.629 -17.057 14.865 1.00 12.59
	ATOM	1417 O LEU 346	3.706 -17.099 14.272 1.00 22.61
	<b>ATOM</b>	1418 N LEU 347	1.612 -17.862 14.574 1.00 18.42
15	ATOM	1419 CA LEU 347	1.706 -18.863 13.517 1.00 18.42
	ATOM	1420 CB LEU 347	0.471 -19.762 13.512 1.00 23.56
	<b>ATOM</b>	1421 CG LEU 347	0.509 -20.965 14.456 1.00 23.56
	ATOM	1422 CD1 LEU 347	-0.819 -21.702 14.398 1.00 23.56
	ATOM	1423 CD2 LEU 347	1.659 -21.890 14.068 1.00 23.56
20	ATOM	1424 C LEU 347	1.870 -18.201 12.154 1.00 18.42
	ATOM	1425 O LEU 347	2.672 -18.651 11.330 1.00 23.56
	ATOM	1426 N ALA 348	1.099 -17.144 11.917 1.00 12.49
	ATOM	1427 CA ALA 348	1.157 -16.403 10.663 1.00 12.49
	ATOM	1428 CB ALA 348	0.098 -15.302 10.654 1.00 14.77
25	ATOM	1429 C ALA 348	2.545 -15.798 10.504 1.00 12.49
	ATOM	1430 O ALA 348	3.154 -15.874 9.436 1.00 14.77
	ATOM	1431 N PHE 349	3.048 -15.246 11.602 1.00 15.52
	ATOM	1432 CA PHE 349	4.357 -14.613 11.664 1.00 15.52
	ATOM	1433 CB PHE 349	4.566 -14.049 13.076 1.00 14.41
30	ATOM	1434 CG PHE 349	5.714 -13.085 13.203 1.00 14.41
	ATOM	1435 CD1 PHE 349	6.473 -12.712 12.099 1.00 14.41
	ATOM	1436 CD2 PHE 349	6.027 -12.540 14.443 1.00 14.41
	ATOM	1437 CE1 PHE 349	7.523 -11.813 12.230 1.00 14.41
2.5	ATOM	1438 CE2 PHE 349	7.075 -11.640 14.584 1.00 14.41
35	ATOM	1439 CZ PHE 349	7.825 -11.275 13.475 1.00 14.41
	ATOM	1440 C PHE 349	5.444 -15.633 11.324 1.00 15.52
	ATOM	1441 O PHE 349	6.252 -15.413 10.422 1.00 14.41
	ATOM	1442 N GLU 350	5.439 -16.760 12.026 1.00 13.20 6.424 -17.811 11.801 1.00 13.20
40	ATOM	1443 CA GLU 350	6.152 -18.995 12.734 1.00 33.43
40	ATOM	1444 CB GLU 350	7.068 -20.193 12.519 1.00 33.43
	ATOM	1445 CG GLU 350	6.786 -21.331 13.482 1.00 33.43
	ATOM	1446 CD GLU 350 1447 OE1 GLU 350	7.746 -22.035 13.857 1.00 33.43
	ATOM		5.611 -21.525 13.865 1.00 33.43
45	ATOM	1448 OE2 GLU 350 1449 C GLU 350	6.409 -18.283 10.352 1.00 13.20
43	ATOM ATOM		7.449 -18.355 9.694 1.00 33.43
	ATOM	1450 O GLU 350 1451 N HIS 351	5.217 -18.573 9.850 1.00 19.10
	ATOM	1451 N HIS 351 1452 CA HIS 351	5.062 -19.051 8.485 1.00 19.10
	ATOM	1452 CA HIS 351 1453 CB HIS 351	3.632 -19.536 8.256 1.00 18.97
50	ATOM	1454 CG HIS 351	3.249 -20.700 9.117 1.00 18.97
50	ATOM	1434 CO 1113 331	3.247 -20.100 3.111 XIOO X0.71

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3.987 -21.474 9.948 1.00 18.97
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                                3.134 -22.394 10.509 1.00 18.97
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     ATOM 1463 CB TYR 352
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     ATOM 1464 CG TYR 352
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     ATOM 1466 CE1 TYR 352
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     ATOM 1468 CE2 TYR 352
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     ATOM 1469 CZ TYR 352
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                                 6.710 -10.401 3.431 1.00 16.06
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     ATOM 1471 C TYR 352
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                                7.851 -15.789 7.980 1.00 15.38
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     ATOM 1476 CG1 VAL 353
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     ATOM 1477 CG2 VAL 353
                                 9.384 -14.703 10.349 1.00 18.40
     ATOM 1478 C VAL 353
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     ATOM 1479 O VAL 353
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     ATOM 1484 OD1 ASN 354
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     ATOM 1485 ND2 ASN 354
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     ATOM 1490 CB HIS 355
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     ATOM 1491 CG HIS 355
     ATOM 1492 CD2 HIS 355
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                               9.834 -16.920 1.595 1.00 24.39
     ATOM 1497 O HIS 355
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     ATOM 1498 N ARG 356
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     ATOM 1499 CA ARG 356
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     ATOM 1500 CB ARG 356
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     ATOM 1501 CG ARG 356
     ATOM 1502 CD ARG 356
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                                10.932 - 10.934 4.592 1.00 22.01
     ATOM 1503 NE ARG 356
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     ATOM 1504 CZ ARG 356
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PCT/US98/25296

#### WO 99/26966

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    ATOM 1506 NH2 ARG 356
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    ATOM 1507 C ARG 356
    ATOM 1508 O ARG 356
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                               12.547 -16.788 3.841 1.00 23.18
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    ATOM 1511 CB LYS 357
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    ATOM 1512 CG LYS 357
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    ATOM 1513 CD LYS 357
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    ATOM 1514 CE LYS 357
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    ATOM 1516 C LYS 357
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    ATOM 1518 N HIS 358
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    ATOM 1519 CA HIS 358
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    ATOM 1520 CB HIS 358
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    ATOM 1521 CG HIS 358
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    ATOM 1522 CD2 HIS 358
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     ATOM 1525 NE2 HIS 358
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     ATOM 1536 CG2 ILE 360
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     ATOM 1537 CG1 ILE 360
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     ATOM 1538 CD1 ILE 360
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     ATOM 1539 C ILE 360
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     ATOM 1545 CG PRO 361
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     ATOM 1546 C PRO 361
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     ATOM 1553 ND1 HIS 362
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     ATOM 1554 CE1 HIS 362
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	ATOM	1555 NE2 HIS 362	20.439 -23.353 14.558 1.00 31.38
	ATOM	1556 C HIS 362	17.189 -18.506 13.628 1.00 18.84
	ATOM	1557 O HIS 362	16.980 -17.979 14.723 1.00 31.38
_	ATOM	1558 N PHE 363	16.825 -17.950 12.476 1.00 18.69
5	ATOM	1559 CA PHE 363	16.209 -16.630 12.408 1.00 18.69
	ATOM	1560 CB PHE 363	15.825 -16.302 10.962 1.00 19.25
	ATOM	1561 CG PHE 363	15.339 -14.894 10.765 1.00 19.25
	ATOM	1562 CD1 PHE 363	16.239 -13.862 10.530 1.00 19.25
	ATOM	1563 CD2 PHE 363	13.981 -14.598 10.819 1.00 19.25
10	ATOM	1564 CE1 PHE 363	15.794 -12.556 10.351 1.00 19.25
	ATOM	1565 CE2 PHE 363	13.527 -13.296 10.642 1.00 19.25
	ATOM	1566 CZ PHE 363	14.435 -12.273 10.407 1.00 19.25
	ATOM	1567 C PHE 363	14.995 -16.461 13.323 1.00 18.69
	ATOM	1568 O PHE 363	14.955 -15.540 14.138 1.00 19.25
15	ATOM	1569 N TRP 364	14.016 -17.351 13.191 1.00 16.46
	ATOM	1570 CA TRP 364	12.797 -17.280 13.995 1.00 16.46
	ATOM	1571 CB TRP 364	11.882 -18.482 13.706 1.00 17.81
	ATOM	1572 CG TRP 364	10.588 -18.488 14.481 1.00 17.81
	ATOM	1573 CD2 TRP 364	9.586 -17.458 14.504 1.00 17.81
20	ATOM	1574 CE2 TRP 364	8.547 -17.905 15.350 1.00 17.81
	ATOM	1575 CE3 TRP 364	9.467 -16.202 13.894 1.00 17.81
	ATOM	1576 CD1 TRP 364	10.126 -19.486 15.290 1.00 17.81
	ATOM	1577 NE1 TRP 364	8.902 -19.144 15.814 1.00 17.81
	ATOM	1578 CZ2 TRP 364	7.403 -17.142 15.602 1.00 17.81
25	ATOM	1579 CZ3 TRP 364	8.329 -15.444 14.145 1.00 17.81
	ATOM	1580 CH2 TRP 364	7.312 -15.919 14.992 1.00 17.81
	ATOM	1581 C TRP 364	13.046 -17.114 15.500 1.00 16.46
	ATOM	1582 O TRP 364	12.595 -16.133 16.087 1.00 17.81
	ATOM	1583 N PRO 365	13.779 -18.051 16.137 1.00 18.31
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	ATOM	1585 CA PRO 365	14.038 -17.920 17.577 1.00 18.31
	ATOM	1586 CB PRO 365	14.939 -19.118 17.874 1.00 25.61
	ATOM	1587 CG PRO 365	14.500 -20.130 16.882 1.00 25.61
	ATOM	1588 C PRO 365	14.732 -16.606 17.933 1.00 18.31
35	ATOM	1589 O PRO 365	14.387 -15.963 18.926 1.00 25.61
	ATOM	1590 N LYS 366	15.699 -16.207 17.112 1.00 25.16
	ATOM	1591 CA LYS 366	16.439 -14.968 17.338 1.00 25.16
	ATOM	1592 CB LYS 366	17.537 -14.805 16.289 1.00 40.51
	ATOM	1593 CG LYS 366	18.679 -15.792 16.417 1.00 40.51
40	ATOM	1594 CD LYS 366	19.664 -15.607 15.278 1.00 40.51
	ATOM	1595 CE LYS 366	20.884 -16.492 15.440 1.00 40.51
	ATOM	1596 NZ LYS 366	21.800 -16.360 14.275 1.00 40.51
	ATOM	1597 C LYS 366	15.521 -13.747 17.317 1.00 25.16
	ATOM	1598 O LYS 366	15.593 -12.893 18.202 1.00 40.51
45	ATOM	1599 N LEU 367	14.661 -13.666 16.307 1.00 25.30
	ATOM	1600 CA LEU 367	13.729 -12.551 16.184 1.00 25.30
	ATOM	1601 CB LEU 367	12.989 -12.620 14.845 1.00 27.80
	ATOM	1602 CG LEU 367	11.964 -11.519 14.561 1.00 27.80
	ATOM	1603 CD1 LEU 367	12.621 -10.147 14.679 1.00 27.80
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     ATOM 1610 CG LEU 368
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    ATOM 1611 CD1 LEU 368
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     ATOM 1612 CD2 LEU 368
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     ATOM 1614 O LEU 368
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    ATOM 1616 CA MET 369
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    ATOM 1617 CB MET 369
     ATOM 1618 CG MET 369
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    ATOM 1619 SD MET 369
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     ATOM 1626 CG LYS 370
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     ATOM 1628 CE LYS 370
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     ATOM 1631 O LYS 370
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     ATOM 1633 CA VAL 371
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     ATOM 1634 CB VAL 371
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     ATOM 1635 CG1 VAL 371
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     ATOM 1636 CG2 VAL 371
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     ATOM 1637 C VAL 371
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     ATOM 1639 N THR 372
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     ATOM 1640 CA THR 372
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     ATOM 1641 CB THR 372
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     ATOM 1642 OG1 THR 372
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     ATOM 1643 CG2 THR 372
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     ATOM 1644 C THR 372
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     ATOM 1645 O THR 372
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     ATOM 1646 N ASP 373
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     ATOM 1647 CA ASP 373
                                14.976 -6.822 23.435 1.00 37.94
     ATOM 1648 CB ASP 373
                                16.065 -7.893 23.445 1.00 37.94
     ATOM 1649 CG ASP 373
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                                 16.248 -8.571 24.483 1.00 37.94
     ATOM 1650 OD1 ASP 373
     ATOM 1651 OD2 ASP 373
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     ATOM 1652 C ASP 373
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                               13.040 -4.928 25.407 1.00 37.94
     ATOM 1653 O ASP 373
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     ATOM 1654 N LEU 374
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     ATOM 1656 CB LEU 374
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     ATOM 1657 CG LEU 374
                                10.624 -4.720 20.985 1.00 20.58
                                 9.826 -5.352 19.862 1.00 20.58
     ATOM 1658 CD1 LEU 374
                                10.599 -3.202 20.882 1.00 20.58
     ATOM 1659 CD2 LEU 374
                               10.145 -4.786 24.825 1.00 17.04
     ATOM 1660 C LEU 374
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     ATOM 1661 O LEU 374
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     ATOM 1664 CB ARG 375
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     ATOM 1665 CG ARG 375
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     ATOM 1666 CD ARG 375
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     ATOM 1667 NE ARG 375
     ATOM 1668 CZ ARG 375
                                5.054 -9.654 25.028 1.00 55.89
                                 4.782 -8.808 24.040 1.00 55.89
     ATOM 1669 NH1 ARG 375
15
     ATOM 1670 NH2 ARG 375
                                 4.438 -10.829 25.073 1.00 55.89
                                9.874 -5.221 27.735 1.00 20.46
     ATOM 1671 C ARG 375
                                9.328 -4.391 28.463 1.00 55.89
     ATOM 1672 O ARG 375
                                11.174 -5.502 27.794 1.00 20.10
     ATOM 1673 N MET 376
                                12.076 -4.863 28.744 1.00 20.10
20
     ATOM 1674 CA MET 376
                                13.493 -5.417 28.580 1.00 63.73
     ATOM 1675 CB MET 376
     ATOM 1676 CG MET 376
                                13.956 -6.310 29.722 1.00 63.73
     ATOM 1677 SD MET 376
                                14.494 -5.373 31.182 1.00 63.73
                                12.934 -5.151 32.087 1.00 63.73
     ATOM 1678 CE MET 376
                               12.081 -3.347 28.566 1.00 20.10
25
     ATOM 1679 C MET 376
                               11.973 -2.602 29.539 1.00 63.73
     ATOM 1680 O MET 376
     ATOM 1681 N ILE 377
                               12.194 -2.896 27.321 1.00 30.02
     ATOM 1682 CA ILE 377
                               12.198 -1.469 27.014 1.00 30.02
     ATOM 1683 CB ILE 377
                               12.329 -1.228 25.488 1.00 19.31
30
     ATOM 1684 CG2 ILE 377
                                12.088 0.242 25.152 1.00 19.31
     ATOM 1685 CG1 ILE 377
                                13.711 -1.685 25.011 1.00 19.31
                                13.906 -1.634 23.507 1.00 19.31
     ATOM 1686 CD1 ILE 377
     ATOM 1687 C ILE 377
                               10.915 -0.821 27.542 1.00 30.02
     ATOM 1688 O ILE 377
                               10.962 0.216 28.211 1.00 19.31
     ATOM 1689 N GLY 378
                                9.779 -1.455 27.266 1.00 21.85
35
     ATOM 1690 CA GLY 378
                                8.505 -0.936 27.729 1.00 21.85
     ATOM 1691 C GLY 378
                                8.459 -0.821 29.243 1.00 21.85
     ATOM 1692 O GLY 378
                                7.990 0.185 29.779 1.00 34.01
                                8.967 -1.842 29.928 1.00 31.30
     ATOM 1693 N ALA 379
                                 8.996 -1.870 31.388 1.00 31.30
     ATOM 1694 CA ALA 379
40
     ATOM 1695 CB ALA 379
                                 9.471 -3.231 31.880 1.00 30.06
     ATOM 1696 C ALA 379
                                9.895 -0.763 31.938 1.00 31.30
     ATOM 1697 O ALA 379
                                9.482 0.002 32.810 1.00 30.06
                               11.117 -0.677 31.418 1.00 28.61
     ATOM 1698 N CYS 380
                                12.067 0.349 31.841 1.00 28.61
     ATOM 1699 CA CYS 380
45
                                13.360 0.268 31.025 1.00 60.26
     ATOM 1700 CB CYS 380
                                14.499 -1.067 31.470 1.00 60.26
     ATOM 1701 SG CYS 380
     ATOM 1702 C CYS 380
                               11.449 1.730 31.658 1.00 28.61
                               11.516 2.573 32.554 1.00 60.26
     ATOM 1703 O CYS 380
                               10.840 1.957 30.498 1.00 30.42
50
     ATOM 1704 N HIS 381
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WO 99/26966

PCT/US98/25296

	ATOM	1705 CA HIS 381	10.212 3.243 30.216 1.00 30.42
	<b>ATOM</b>	1706 CB HIS 381	9.696 3.306 28.779 1.00 16.49
	<b>ATOM</b>	1707 CG HIS 381	8.942 4.562 28.472 1.00 16.49
	ATOM	1708 CD2 HIS 381	9.370 5.805 28.151 1.00 16.49
5	ATOM	1709 ND1 HIS 381	7.566 4.633 28.524 1.00 16.49
	ATOM	1710 CE1 HIS 381	7.180 5.866 28.251 1.00 16.49
	<b>ATOM</b>	1711 NE2 HIS 381	8.255 6.596 28.021 1.00 16.49
	ATOM	1712 C HIS 381	9.073 3.539 31.182 1.00 30.42
	4 TO 1 4	1712 O LIC 201	0 056 4 600 21 552 1 00 16 40

ATOM 1713 O HIS 381 8.856 4.690 31.552 1.00 16.49 8.330 2.506 31.564 1.00 22.89 10 ATOM 1714 N ALA 382 7.218 2.666 32.493 1.00 22.89 ATOM 1715 CA ALA 382 6.520 1.336 32.708 1.00 34.50 ATOM 1716 CB ALA 382 7.738 3.213 33.819 1.00 22.89 ATOM 1717 C ALA 382 ATOM 1718 O ALA 382 7.219 4.200 34.343 1.00 34.50 ATOM 1719 N SER 383 8.789 2.586 34.336 1.00 26.39 15

ATOM 1720 CA SER 383 9.400 3.006 35.591 1.00 26.39 ATOM 1721 CB SER 383 10.510 2.030 35.985 1.00 52.94 ATOM 1722 OG SER 383 10.015 0.702 36.046 1.00 52.94 ATOM 1723 C SER 383 9.966 4.418 35.470 1.00 26.39 20 ATOM 1724 O SER 383 9.772 5.253 36.357 1.00 52.94

ATOM 1724 O SER 383 9.772 5.253 36.357 1.00 32.94 ATOM 1725 N ARG 384 10.662 4.683 34.368 1.00 30.36 ATOM 1726 CA ARG 384 11.249 5.995 34.134 1.00 30.36 ATOM 1727 CB ARG 384 12.116 5.977 32.874 1.00 37.39 ATOM 1728 CG ARG 384 12.601 7.344 32.431 1.00 37.39 ATOM 1729 CD ARG 384 14.070 7.321 32.060 1.00 37.39 ATOM 1730 NE ARG 384 14.935 7.597 33.204 1.00 37.39

ATOM 1731 CZ ARG 384 15.750 8.646 33.291 1.00 37.39 ATOM 1732 NH1 ARG 384 15.824 9.529 32.303 1.00 37.39 ATOM 1733 NH2 ARG 384 16.488 8.819 34.376 1.00 37.39 ATOM 1734 C ARG 384 10.169 7.067 34.030 1.00 30.36 ATOM 1735 O ARG 384 10.301 8.144 34.616 1.00 37.39

ATOM 1736 N PHE 385 9.078 6.749 33.338 1.00 24.47 ATOM 1737 CA PHE 385 7.980 7.693 33.171 1.00 24.47 ATOM 1738 CB PHE 385 6.859 7.092 32.319 1.00 28.70 ATOM 1740 CD1 PHE 385 5.710 8.036 32.075 1.00 28.70 ATOM 1741 CD2 PHE 385 5.795 9.017 31.092 1.00 28.70 ATOM 1742 CE1 PHE 385 4.549 7.954 32.836 1.00 28.70 ATOM 1742 CE1 PHE 385 4.740 9.903 30.874 1.00 28.70

ATOM 1743 CE2 PHE 385 3.491 8.835 32.624 1.00 28.70 40 ATOM 1744 CZ PHE 385 3.587 9.812 31.641 1.00 28.70 ATOM 1745 C PHE 385 7.436 8.097 34.533 1.00 24.47

ATOM 1746 O PHE 385 7.250 9.285 34.805 1.00 28.70 ATOM 1747 N LEU 386 7.208 7.107 35.391 1.00 31.13 ATOM 1748 CA LEU 386 6.690 7.352 36.734 1.00 31.13 ATOM 1749 CB LEU 386 6.596 6.044 37.513 1.00 39.10

45 ATOM 1749 CB LEU 386 6.596 6.044 37.513 1.00 39.10 ATOM 1750 C LEU 386 7.577 8.348 37.474 1.00 31.13 ATOM 1751 O LEU 386 7.085 9.201 38.217 1.00 39.10 ATOM 1752 N HIS 387 8.884 8.254 37.243 1.00 36.46 ATOM 1753 CA HIS 387 9.837 9.152 37.881 1.00 36.46

50 ATOM 1754 CB HIS 387 9.637 9.132 37.881 1.00 50.40

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11.459 7.338 38.590 1.00 62.78
    ATOM 1755 CG HIS 387
                                10.601 6.614 39.346 1.00 62.78
     ATOM 1756 CD2 HIS 387
                                12.675 6.689 38.663 1.00 62.78
    ATOM 1757 ND1 HIS 387
                               12.554 5.620 39.431 1.00 62.78
    ATOM 1758 CE1 HIS 387
                               11.309 5.550 39.856 1.00 62.78
    ATOM 1759 NE2 HIS 387
                               9.778 10.544 37.266 1.00 36.46
     ATOM 1760 C HIS 387
                               9.885 11.543 37.979 1.00 62.78
    ATOM 1761 O HIS 387
                                9.587 10.612 35.950 1.00 33.41
     ATOM 1762 N MET 388
    ATOM 1763 CA MET 388
ATOM 1764 CB MET 388
                                 9.505 11.894 35.258 1.00 33.41
                                9.269 11.703 33.755 1.00 42.63
10
                                10.456 11.144 32.982 1.00 42.63
    ATOM 1765 CG MET 388
                                10.253 11.325 31.192 1.00 42.63
    ATOM 1766 SD MET 388
                                9.501 9.772 30.748 1.00 42.63
    ATOM 1767 CE MET 388
    ATOM 1768 C MET 388
                                8.385 12.746 35.849 1.00 33.41
     ATOM 1769 O MET 388
                                8.573 13.934 36.103 1.00 42.63
15
                               7.235 12.126 36.092 1.00 39.26
     ATOM 1770 N LYS 389
                                6.082 12.825 36.659 1.00 39.26
    ATOM 1771 CA LYS 389
                                4.867 11.900 36.719 1.00 52.87
    ATOM 1772 CB LYS 389
                                4.237 11.594 35.379 1.00 52.87
     ATOM 1773 CG LYS 389
20
    ATOM 1774 CD LYS 389
                                3.048 10.667 35.553 1.00 52.87
                                3.482 9.327 36.125 1.00 52.87
     ATOM 1775 CE LYS 389
                                2.335 8.407 36.326 1.00 52.87
     ATOM 1776 NZ LYS 389
                               6.363 13.360 38.056 1.00 39.26
     ATOM 1777 C LYS 389
     ATOM 1778 O LYS 389
                               5.837 14.404 38.452 1.00 52.87
     ATOM 1779 N VAL 390
                                7.156 12.614 38.818 1.00 44.18
25
     ATOM 1780 CA VAL 390
                                7.508 13.016 40.172 1.00 44.18
                                8.299 11.898 40.905 1.00 50.50
     ATOM 1781 CB VAL 390
     ATOM 1782 CG1 VAL 390
                                 8.718 12.362 42.293 1.00 50.50
     ATOM 1783 CG2 VAL 390
                                 7.455 10.640 41.012 1.00 50.50
     ATOM 1784 C VAL 390
                                8.352 14.288 40.145 1.00 44.18
30
     ATOM 1785 O VAL 390
                                8.144 15.198 40.948 1.00 50.50
                                9.261 14.368 39.179 1.00 38.64
     ATOM 1786 N GLU 391
                                10.161 15.509 39.056 1.00 38.64
     ATOM 1787 CA GLU 391
     ATOM 1788 CB GLU 391
                                11.483 15.060 38.424 1.00 64.18
     ATOM 1789 CG GLU 391
                                12.065 13.766 39.009 1.00 64.18
35
    ATOM 1790 CD GLU 391
                                12.662 13.922 40.405 1.00 64.18
     ATOM 1791 OE1 GLU 391
                                12.190 14.773 41.192 1.00 64.18
                                13.611 13.173 40.721 1.00 64.18
     ATOM 1792 OE2 GLU 391
     ATOM 1793 C GLU 391
                                9.623 16.737 38.314 1.00 38.64
     ATOM 1794 O GLU 391
                                9.656 17.850 38.849 1.00 64.18
40
                                9.125 16.539 37.096 1.00 37.24
     ATOM 1795 N CYS 392
                                8.611 17.635 36.271 1.00 37.24
     ATOM 1796 CA CYS 392
                                8.879 17.345 34.784 1.00 30.64
     ATOM 1797 CB CYS 392
                                10.634 17.137 34.283 1.00 30.64
     ATOM 1798 SG CYS 392
     ATOM 1799 C CYS 392
                                7.110 17.882 36.496 1.00 37.24
45
                                6.403 17.011 37.006 1.00 30.64
     ATOM 1800 O CYS 392
                                6.625 19.107 36.199 1.00 40.56
     ATOM 1801 N PRO 393
     ATOM 1802 CD PRO 393
                                7.444 20.297 35.904 1.00 33.41
                                5.209 19.473 36.358 1.00 40.56
     ATOM 1803 CA PRO 393
                                5.253 21.001 36.404 1.00 33.41
     ATOM 1804 CB PRO 393
50
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	ATOM	1805 CG PRO 393	6.409 21.332 35.527 1.00 33.41
	ATOM	1806 C PRO 393	4.330 18.975 35.207 1.00 40.56
		1807 O PRO 393	4.776 18.907 34.057 1.00 33.41
	ATOM	1808 N THR 394	3.067 18.691 35.516 1.00 41.91
-	ATOM		2.101 18.186 34.540 1.00 41.91
5	ATOM	1809 CA THR 394 1810 CB THR 394	
	ATOM	= = : :	0.691 18.075 35.156 1.00 62.04 0.706 18.582 36.497 1.00 62.04
	ATOM		0.706 18.382 36.497 1.00 62.04
	ATOM		
10	ATOM	1813 C THR 394	1.995 18.984 33.242 1.00 41.91 1.758 18.411 32.181 1.00 62.04
10	ATOM	1814 O THR 394	2.191 20.297 33.327 1.00 43.92
	ATOM	1815 N GLU 395	2.191 20.297 33.327 1.00 43.92 2.104 21.176 32.160 1.00 43.92
	ATOM	1816 CA GLU 395	2.104 21.176 32.160 1.00 43.92 2.313 22.626 32.585 1.00 34.22
	ATOM	1817 CB GLU 395	
	ATOM	1818 C GLU 395	3.071 20.814 31.031 1.00 43.92
15	ATOM	1819 O GLU 395	2.887 21.243 29.891 1.00 34.22
	ATOM	1820 N LEU 396	4.104 20.041 31.350 1.00 34.92
	ATOM	1821 CA LEU 396	5.096 19.634 30.359 1.00 34.92
	ATOM	1822 CB LEU 396	6.473 19.495 31.017 1.00 35.81
•	ATOM	1823 CG LEU 396	7.074 20.747 31.662 1.00 35.81
20	ATOM	1824 CD1 LEU 396	8.427 20.410 32.263 1.00 35.81
	ATOM	1825 CD2 LEU 396	7.209 21.857 30.629 1.00 35.81
	ATOM	1826 C LEU 396	4.731 18.324 29.661 1.00 34.92
	ATOM	1827 O LEU 396	5.343 17.954 28.659 1.00 35.81
	ATOM	1828 N PHE 397	3.734 17.627 30.197 1.00 35.28
25	ATOM	1829 CA PHE 397	3.302 16.352 29.640 1.00 35.28
	ATOM	1830 CB PHE 397	3.059 15.341 30.764 1.00 27.13
	ATOM	1831 CG PHE 397	4.285 15.004 31.561 1.00 27.13
	ATOM	1832 CD1 PHE 397	4.700 15.824 32.604 1.00 27.13 5.021 13.860 31.273 1.00 27.13
••	ATOM	1833 CD2 PHE 397	
30	ATOM	1834 CE1 PHE 397	5.831 15.510 33.349 1.00 27.13
	ATOM	1835 CE2 PHE 397	6.155 13.537 32.013 1.00 27.13
	ATOM	1836 CZ PHE 397	6.561 14.364 33.052 1.00 27.13
	ATOM	1837 C PHE 397	2.027 16.474 28.812 1.00 35.28
	ATOM	1838 O PHE 397	0.977 16.861 29.331 1.00 27.13
35	ATOM	1839 N PRO 398	2.102 16.164 27.505 1.00 26.41
	ATOM	1840 CD PRO 398	3.305 15.850 26.713 1.00 19.32
	ATOM	1841 CA PRO 398	0.917 16.247 26.647 1.00 26.41
	ATOM	1842 CB PRO 398	1.439 15.752 25.300 1.00 19.32
	ATOM	1843 CG PRO 398	2.867 16.193 25.312 1.00 19.32
40	ATOM	1844 C PRO 398	-0.157 15.313 27.206 1.00 26.41
	ATOM	1845 O PRO 398	0.160 14.232 27.710 1.00 19.32
	ATOM	1846 N PRO 399	-1.439 15.702 27.104 1.00 25.12
	ATOM	1847 CD PRO 399	-1.935 16.929 26.454 1.00 24.32
	ATOM	1848 CA PRO 399	-2.554 14.894 27.612 1.00 25.12
45	ATOM	1849 CB PRO 399	-3.777 15.594 27.022 1.00 24.32
	ATOM	1850 CG PRO 399	-3.349 17.026 26.974 1.00 24.32
	ATOM	1851 C PRO 399	-2.502 13.416 27.222 1.00 25.12
	ATOM	1852 O PRO 399	-2.599 12.540 28.085 1.00 24.32
	ATOM	1853 N LEU 400	-2.322 13.139 25.933 1.00 23.10
50	ATOM	1854 CA LEU 400	-2.265 11.759 25.454 1.00 23.10

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-2.230 11.720 23.923 1.00 22.35
    ATOM 1855 CB LEU 400
                               -2.485 10.354 23.276 1.00 22.35
    ATOM 1856 CG LEU 400
                                -3.792 9.765 23.792 1.00 22.35
    ATOM 1857 CD1 LEU 400
    ATOM 1858 CD2 LEU 400
                                -2.523 10,494 21.763 1.00 22.35
                              -1.066 11.012 26.032 1.00 23.10
    ATOM 1859 C LEU 400
    ATOM 1860 O LEU 400
                               -1.160 9.825 26.345 1.00 22.35
                               0.044 11.723 26.202 1.00 13.85
    ATOM 1861 N PHE 401
    ATOM 1862 CA PHE 401
                               1.269 11.150 26.755 1.00 13.85
                                2.374 12.213 26.753 1.00 26.97
    ATOM 1863 CB PHE 401
                                3.729 11.702 27.164 1.00 26.97
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    ATOM 1864 CG PHE 401
                                4.189 10.461 26.732 1.00 26.97
    ATOM 1865 CD1 PHE 401
                                4.561 12.481 27.963 1.00 26.97
    ATOM 1866 CD2 PHE 401
    ATOM 1867 CE1 PHE 401
                                5.459 10.005 27.091 1.00 26.97
    ATOM 1868 CE2 PHE 401
                                5.830 12.035 28.327 1.00 26.97
                                6.280 10.795 27.889 1.00 26.97
    ATOM 1869 CZ PHE 401
15
    ATOM 1870 C PHE 401
                               0.993 10.659 28.179 1.00 13.85
    ATOM 1871 O PHE 401
                               1.393 9.558 28.555 1.00 26.97
                               0.274 11.473 28.947 1.00 25.21
    ATOM 1872 N LEU 402
    ATOM 1873 CA LEU 402
                               -0.080 11.145 30.325 1.00 25.21
    ATOM 1874 CB LEU 402
                               -0.640 12.380 31.035 1.00 29.34
20
    ATOM 1875 CG LEU 402
                                0.334 13.411 31.600 1.00 29.34
                                -0.430 14.658 32.018 1.00 29.34
    ATOM 1876 CD1 LEU 402
    ATOM 1877 CD2 LEU 402
                                1.090 12.814 32.775 1.00 29.34
                               -1.109 10.025 30.425 1.00 25.21
    ATOM 1878 C LEU 402
                               -1.034 9.189 31.320 1.00 29.34
    ATOM 1879 O LEU 402
25
                               -2.090 10.043 29.529 1.00 23.54
    ATOM 1880 N GLU 403
                               -3.159 9.046 29.521 1.00 23.54
    ATOM 1881 CA GLU 403
                                -4.274 9.482 28.562 1.00 63.22
    ATOM 1882 CB GLU 403
                                -5.469 8.531 28.506 1.00 63.22
    ATOM 1883 CG GLU 403
                                -6.530 8.952 27.498 1.00 63.22
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    ATOM 1884 CD GLU 403
    ATOM 1885 OE1 GLU 403
                                -6.237 9.786 26.613 1.00 63.22
                                -7.666 8.436 27.589 1.00 63.22
    ATOM 1886 OE2 GLU 403
                               -2.708 7.629 29.170 1.00 23.54
    ATOM 1887 C GLU 403
    ATOM 1888 O GLU 403
                               -3.210 6.656 29.735 1.00 63.22
                               -1.787 7.515 28.221 1.00 33.24
    ATOM 1889 N VAL 404
35
                                -1.297 6.213 27.782 1.00 33.24
    ATOM 1890 CA VAL 404
                                -0.621 6.314 26.390 1.00 30.71
    ATOM 1891 CB VAL 404
                                -0.097 4.957 25.947 1.00 30.71
    ATOM 1892 CG1 VAL 404
                                -1.611 6.841 25.371 1.00 30.71
    ATOM 1893 CG2 VAL 404
                               -0.338 5.528 28.752 1.00 33.24
40
    ATOM 1894 C VAL 404
                               -0.386 4.305 28.914 1.00 30.71
    ATOM 1895 O VAL 404
                               0.526 6.309 29.392 1.00 33.66
     ATOM 1896 N PHE 405
     ATOM 1897 CA PHE 405
                                1.516 5.752 30.308 1.00 33.66
     ATOM 1898 CB PHE 405
                                2.901 6.326 29.984 1.00 34.35
     ATOM 1899 CG PHE 405
                                3.343 6.076 28.568 1.00 34.35
45
                                 3.519 7.134 27.683 1.00 34.35
     ATOM 1900 CD1 PHE 405
                                 3.569 4.782 28.114 1.00 34.35
     ATOM 1901 CD2 PHE 405
                                3.911 6.906 26.365 1.00 34.35
     ATOM 1902 CE1 PHE 405
                                3.960 4.545 26.798 1.00 34.35
     ATOM 1903 CE2 PHE 405
                                4.131 5.610 25.922 1.00 34.35
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    ATOM 1904 CZ PHE 405
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ATOM 1905 C PHE 405
                               1.189 5.931 31.790 1.00 33.66
     ATOM 1906 O PHE 405
                               2.036 5.539 32.623 1.00 34.35
     ATOM 1907 OXT PHE 405
                                 0.090 6.434 32.107 1.00 34.35
     ATOM 1908 C1 TRI
                              8.375 7.063 18.475 1.00 34.21
                          1
    ATOM 1909 C2 TRI
                              10.048 8.688 23.016 1.00 33.36
                          1
     ATOM 1910 C3 TRI
                              8.104 8.391 18.941 1.00 34.21
                         1
                              10.496 9.696 23.813 1.00 33.36
    ATOM 1911 C4 TRI
                          1
                              8.916 8.943 19.927 1.00 34.21
    ATOM 1912 C5 TRI
                          1
                            10.152 9.772 25.121 1.00 33.36
    ATOM 1913 C6 TRI
                          1
    ATOM 1914 C7 TRI
                            9.862 8.178 20.609 1.00 34.21
10
                          1
                              9.246 8.821 25.653 1.00 33.36
    ATOM 1915 C8 TRI
                         1
                            10.117 6.865 20.147 1.00 34.21
    ATOM 1916 C9 TRI
                         1
                            8.805 7.754 24.847 1.00 33.36
    ATOM 1917 C10 TRI
                         1
                              9.375 6.339 19.026 1.00 34.21
    ATOM 1918 C11 TRI
                         1
                              9.125 7.756 23.490 1.00 33.36
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     ATOM 1919 C12 TRI
                          1
     ATOM 1920 C13 TRI
                              7.540 6.470 17.383 1.00 35.85
                          1
    ATOM 1921 C15 TRI
                              8.158 6.555 15.938 1.00 35.85
                          1
                             8.713 10.990 20.395 1.00 34.21
    ATOM 1922 I1 TRI
                        1
     ATOM 1923 I2 TRI
                             10.951 11.289 26.315 1.00 33.36
                         1
     ATOM 1924 I3 TRI
                             11.592 5.685 21.118 1.00 34.21
20
                         1
    ATOM 1925 O3 TRI
                             9.407 6.654 15.852 1.00 35.85
                         1
    ATOM 1926 O2 TRI
                            10.570 8.649 21.717 1.00 33.36
                          1
     ATOM 1927 O1 TRI
                              8.798 8.969 26.979 1.00 33.36
                          1
     ATOM 1928 O4 TRI
                              7.352 6.522 14.973 1.00 35.85
                          1
    ATOM 1929 O1 HOH 501
                                9.189 2.098 11.091 1.00 33.36
25
    ATOM 1930 O1 HOH 503
                                5.152 5.261 12.137 1.00 33.36
                                3.970 5.057 16.390 1.00 33.36
    ATOM 1931 O1 HOH 504
    ATOM 1932 O1 HOH 534
                                8.296 -0.941 8.998 1.00 33.36
                                4.845 14.369 13.635 1.00 33.36
     ATOM 1933 O1 HOH 538
     ATOM 1934 O1 HOH 540
                                5.789 12.049 10.352 1.00 33.36
30
                                5.721 2.525 28.939 1.00 33.36
    ATOM 1936 O1 HOH 555
    ATOM 1937 O1 HOH 556
                                3.732 1.273 26.724 1.00 33.36
    ATOM 1935 O1 HOH 600
                                8.767 4.847 8.517 1.00 33.36
     ATOM 1938 AS1 CAD 701
                                1.863 1.579 0.837 1.00 37.00
35
     ATOM 1939 C2 CAD 701
                                1.760 -0.100 0.335 1.00 33.36
     ATOM 1940 C3 CAD 701
                                3.511 1.872 1.858 1.00 28.02
                                1.785 2.506 -0.433 1.00 28.02
    ATOM 1941 O4 CAD 701
                                0.592 2.019 1.654 1.00 28.02
    ATOM 1942 O5 CAD 701
    ATOM 1943 AS AS 801
                              11.254 16.718 33.126 1.00 37.00
                                                             AS
                               16.338 -1.161 29.914 1.00 37.00
                                                             AS
     ATOM 1944 AS AS
                         802
40
     ATOM 1945 AS AS 803
                             -14.931 -11.763 25.324 1.00 37.00
                                                             AS
    END
```

WO 99/26966

WO 99/26966 PCT/US98/25296

#### APPENDIX 5

#### TR\_IPBR2.PDB

REMARK rTR\_ipbr2 full length numbering

REMARK

5 REMARK Rfactor 0.214 Rfree 0.224

REMARK Resolution 15. 2.2 all reflections

REMARK

REMARK Three cacodylate-modified cysteines (CYA)

REMARK Cya334, Cya380, Cya392

10 REMARK cacodylate modeled as single arsenic atom

**REMARK** 

REMARK side chain of certain residues modeled as ALA due to poor

density;

REMARK however, residue name reflects true residue for clarity

15 REMARK

REMARK clone obtained from Murray et. al.

REMARK deposited sequence confirmed,

REMARK differing from that reported by Thompson et. al.

REMARK in the following codons:

20 REMARK 281 Thr - Ala

REMARK 285 Lys - Glu

REMARK identical to that reported by Mitsuhashi et. al.

REMARK gb:RNTRAVI X07409

JRNL AUTH M.B. MURRAY, N.D.ZILZ,

25 N.L.MCCREARY, M.J.MACDONALD

JRNL AUTH 2 H.C.TOWLE

JRNL TITL ISOLATION AND CHARACTERIZATION OF RAT CDNA CLONES FOR

TWO

30 JRNL TITL 2 DISTINCT THYROID HORMONE RECPTORS

JRNL REF JBC

JRNL AUTH C.C.THOMPSON, C.WEINBERGER, R.LEBO, R.M.EVANS

V. 263 25 1988

JRNL TITL IDENTIFICATION OF A NOVEL THYROID HORMONE RECEPTOR

35 EXPRESSED

JRNL TITL 2 IN THE MAMMALIAN CENTRAL NERVOUS SYSTEM

JRNL REF SCIENCE

JRNL AUTH T.MITSUHASHI,G.TENNYSON,V.NIKODEM

JRNL TITL NUCLEOTIDE SEQUENCE OF NOVEL CDNAS GENERATED

V. 237 1987

40 BY

**ALTERNATIVE** 

JRNL TITL 2 SPLICING OF A RAT THYROID HORMONE RECEPTOR

GENE

TRANSCRIPT

45 JRNL REF NUC. ACIDS. RES. V. 16 12 1988

**REMARK** 

ATOM 1 CB ARG 157 68.481 10.663 6.906 1.00 57.50

ATOM 2 CG ARG 157 69.793 10.213 7.512 1.00 59.93

	ATOM	3 CD ARG 157	70.510 11.365 8.189 1.00 70.24
		4 NE ARG 157	71.661 10.906 8.961 1.00 77.62
	ATOM	5 CZ ARG 157	71.599 10.492 10.224 1.00 78.75
	ATOM	6 NH1 ARG 157	70.440 10.480 10.870 1.00 74.33
_	ATOM		72.697 10.075 10.839 1.00 83.44
5	ATOM		66.314 10.014 5.809 1.00 46.84
	ATOM		66.109 10.397 4.659 1.00 54.49
	ATOM		68.442 9.069 5.013 1.00 56.54
	ATOM		67.704 9.537 6.222 1.00 52.92
10	ATOM		65.335 9.953 6.727 1.00 39.44
10	ATOM		
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	ATOM	14 CA PRO 158	63.946 10.368 6.487 1.00 34.98
	ATOM	15 CB PRO 158	63.282 10.172 7.854 1.00 34.92
	ATOM	16 CG PRO 158	64.096 9.096 8.487 1.00 45.83
15	ATOM	17 C PRO 158	63.765 11.804 5.992 1.00 34.13
	ATOM	18 O PRO 158	64.223 12.757 6.621 1.00 31.07
	ATOM	19 N GLU 159	63.110 11.932 4.841 1.00 31.36
	ATOM	20 CA GLU 159	62.814 13.220 4.228 1.00 27.34
	ATOM	21 CB GLU 159	62.569 13.041 2.726 1.00 24.27
20	ATOM	22 CG GLU 159	63.814 12.866 1.887 1.00 24.85
	ATOM	23 CD GLU 159	64.409 14.188 1.454 1.00 28.12
	ATOM	24 OE1 GLU 159	63.642 15.144 1.224 1.00 29.26
	ATOM	25 OE2 GLU 159	65.646 14.269 1.326 1.00 29.52
	ATOM	26 C GLU 159	61.528 13.707 4.870 1.00 24.30
25	ATOM	27 O GLU 159	60.855 12.934 5.566 1.00 29.01
	ATOM	28 N PRO 160	61.192 14.989 4.718 1.00 24.62
	ATOM	29 CD PRO 160	61.979 16.126 4.188 1.00 18.72
	ATOM	30 CA PRO 160	59.947 15.451 5.330 1.00 21.62
	ATOM	31 CB PRO 160	59.945 16.955 5.048 1.00 12.71
30	ATOM	32 CG PRO 160	61.394 17.297 4.930 1.00 15.12
	ATOM	33 C PRO 160	58.743 14.752 4.671 1.00 24.61
	ATOM	34 O PRO 160	58.789 14.384 3.490 1.00 22.63
	ATOM	35 N THR 161	57.705 14.504 5.450 1.00 25.86
	ATOM	36 CA THR 161	56.515 13.864 4.921 1.00 23.77
35	ATOM	37 CB THR 161	55.689 13.201 6.048 1.00 21.75
	ATOM	38 OG1 THR 161	55.178 14.210 6.926 1.00 20.78
	ATOM	39 CG2 THR 161	56.549 12.227 6.847 1.00 18.44
	ATOM	40 C THR 161	55.680 14.967 4.269 1.00 28.67
	ATOM	41 O THR 161	55.917 16.151 4.510 1.00 29.90
40	ATOM	42 N PRO 162	54.685 14.597 3.448 1.00 27.79
	ATOM	43 CD PRO 162	54.313 13.237 3.019 1.00 23.25
	ATOM	44 CA PRO 162	53.843 15.603 2.795 1.00 26.19
	ATOM	45 CB PRO 162	52.699 14.766 2.227 1.00 19.89
	ATOM	46 CG PRO 162	53.394 13.492 1.848 1.00 20.63
45	ATOM	47 C PRO 162	53.334 16.661 3.775 1.00 24.81
	<b>ATOM</b>	48 O PRO 162	53.477 17.863 3.526 1.00 21.10
	ATOM	49 N GLU 163	52.812 16.198 4.911 1.00 26.34
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	<b>ATOM</b>	51 CB GLU 163	51.640 16.231 7.086 1.00 29.46
50	ATOM	52 CG GLU 163	50.482 15.321 6.666 1.00 48.37

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             84 CG ASP 166
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                                51.816 21.028 2.360 1.00 26.38
             85 OD1 ASP 166
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     ATOM
             86 OD2 ASP 166
                                50.464 19.803 3.570 1.00 32.25
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             94 CD2 LEU 167
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                               55.501 23.533 7.904 1.00 23.19
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                                59.740 23.451 5.304 1.00 16.71
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             101 CG1 ILE 168
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             102 CD1 ILE 168
                                60.063 20.588 6.264 1.00 18.18
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Page 236 of 447

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                               54,638 26.782 5.745 1.00 19.88
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                                52.755 27.100 7.330 1.00 26.06
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            117 CB VAL 170
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                               56.877 34.141 6.498 1.00 27.76
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                               56.071 32.383 7.664 1.00 25.31
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             145 CB ALA 174
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                               59.503 33.769 9.592 1.00 20.21
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             150 CB HIS 175
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             152 CD2 HIS 175
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                                66.572 43.877 6.552 1.00 37.42
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            199 CD GLN 181
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WO 99/26966

PCT/US98/25296

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		204 N GL1 182 205 CA GLY 182	65,074 39.713 4.732 1.00 46.26
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-	ATOM		67.309 39.175 5.419 1.00 56.26
5	ATOM	207 O GLY 182	
	ATOM	208 N SER 183	
	ATOM	209 CA SER 183	•
	ATOM	210 CB SER 183	
	ATOM	211 OG SER 183	
10	ATOM	212 C SER 183	69.121 40.197 2.558 1.00 59.84
	ATOM	213 O SER 183	70.352 40.138 2.540 1.00 66.02
	ATOM	214 N HIS 184	68.453 41.338 2.413 1.00 60.68
	ATOM	215 CA HIS 184	69.131 42.600 2.139 1.00 60.01
	ATOM	216 CB HIS 184	68.150 43.596 1.517 1.00 53.49
15	ATOM	217 C HIS 184	69.798 43.209 3.380 1.00 59.43
	ATOM	218 O HIS 184	70.373 44.300 3.303 1.00 59.56
	ATOM	219 N TRP 185	69.753 42.500 4.508 1.00 57.54
	ATOM	220 CA TRP 185	70.343 42.995 5.754 1.00 54.25
•	ATOM	221 CB TRP 185	70.147 41.988 6.899 1.00 47.54
20	ATOM	222 CG TRP 185	70.905 40.692 6.752 1.00 41.08
	ATOM	223 CD2 TRP 185	72.233 40.404 7.230 1.00 39.59
	ATOM	224 CE2 TRP 185	72.522 39.070 6.874 1.00 30.27
	ATOM	225 CE3 TRP 185	73.202 41.146 7.919 1.00 35.23
25	ATOM	226 CD1 TRP 185	70.462 39.553 6.149 1.00 39.73 71.427 38.577 6.219 1.00 40.01
25	ATOM	227 NE1 TRP 185	
	ATOM	228 CZ2 TRP 185	
	ATOM	229 CZ3 TRP 185	
	ATOM	230 CH2 TRP 185 231 C TRP 185	74.673 39.203 7.861 1.00 31.71 71.818 43.382 5.655 1.00 54.21
20	ATOM		
30	ATOM ATOM	232 O TRP 185 233 N LYS 186	72.229 44.403 6.200 1.00 52.82 72.605 42.584 4.938 1.00 54.57
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			74.712 41.682 4.080 1.00 53.31
	ATOM ATOM	235 CB LYS 186 236 C LYS 186	74.712 41.082 4.080 1.00 53.51
35	ATOM	237 O LYS 186	75.417 44.731 4.226 1.00 62.57
33	ATOM	237 O L13 180 238 N GLN 187	73.382 44.640 3.268 1.00 60.12
	ATOM	239 CA GLN 187	73.563 45.873 2.512 1.00 60.15
	ATOM	240 CB GLN 187	73.157 45.653 1.050 1.00 57.00
	ATOM	241 C GLN 187	72.809 47.064 3.101 1.00 60.91
40	ATOM	241 C GLN 187 242 O GLN 187	73.149 48.213 2.822 1.00 66.50
40	ATOM	242 O GLN 187 243 N ARG 188	71.795 46.790 3.919 1.00 59.55
	ATOM	244 CA ARG 188	70.983 47.847 4.525 1.00 59.26
	ATOM	245 CB ARG 188	69.504 47.462 4.466 1.00 55.21
	ATOM	246 C ARG 188	71.372 48.243 5.959 1.00 58.97
45	ATOM	247 O ARG 188	70.914 49.269 6.469 1.00 58.54
47	ATOM	248 N ARG 189	72.202 47.432 6.607 1.00 55.46
	ATOM	249 CA ARG 189	72.630 47.704 7.979 1.00 52.98
	ATOM	250 CB ARG 189	73.211 46.437 8.619 1.00 47.73
	ATOM	251 CG ARG 189	74.509 45.985 7.989 1.00 47.88
50	ATOM	251 CG ARG 189 252 CD ARG 189	75.080 44.763 8.654 1.00 46.96
50	AIOWI	232 CD ARG 107	75.000 14.705 0.051 1100 10.70

PCT/US98/25296

### WO 99/26966

	ATOM	253 NE ARG 189	76.377 44.441 8.068 1.00 57.93
	ATOM	254 CZ ARG 189	77.450 44.090 8.768 1.00 64.81
	ATOM	255 NH1 ARG 189	77.385 44.005 10.087 1.00 67.27
	<b>ATOM</b>	256 NH2 ARG 189	78.600 43.860 8.148 1.00 67.84
5	ATOM	257 C ARG 189	73.650 48.838 8.091 1.00 53.48
	ATOM	258 O ARG 189	74.513 49.004 7.227 1.00 57.14
	<b>ATOM</b>	259 N LYS 190	73.533 49.617 9.161 1.00 51.31
	ATOM	260 CA LYS 190	74.444 50.722 9.435 1.00 48.83
	ATOM	261 CB LYS 190	73.682 52.036 9.516 1.00 45.36
10	ATOM	262 C LYS 190	75.101 50.411 10.773 1.00 46.88
	ATOM	263 O LYS 190	74.454 49.872 11.675 1.00 48.81
	ATOM	264 N PHE 191	76.385 50.724 10.894 1.00 46.98
	ATOM	265 CA PHE 191	77.123 50.462 12.125 1.00 44.38
	ATOM	266 CB PHE 191	78.630 50.520 11.873 1.00 44.25
15	ATOM	267 CG PHE 191	79.170 49.336 11.123 1.00 49.51
13	ATOM	268 CD1 PHE 191	78.828 49.124 9.791 1.00 52.20
	ATOM	269 CD2 PHE 191	80.029 48.437 11.748 1.00 47.25
	ATOM	270 CE1 PHE 191	79.335 48.031 9.090 1.00 55.86
	ATOM	270 CETTHE 191 271 CE2 PHE 191	80.542 47.343 11.059 1.00 49.73
20	ATOM	271 CEZ PHE 191	80.195 47.139 9.727 1.00 51.55
20	ATOM	273 C PHE 191	76.764 51.443 13.233 1.00 46.44
	ATOM	274 O PHE 191	76.647 52.645 12.996 1.00 51.28
	ATOM	274 O FHE 191 275 N LEU 192	76.567 50.924 14.439 1.00 47.66
	ATOM	276 CA LEU 192	76.256 51.776 15.577 1.00 46.44
25	_	277 CB LEU 192	75.930 50.924 16.808 1.00 38.06
25	ATOM		75.527 51.672 18.082 1.00 33.55
	ATOM		74.180 52.339 17.871 1.00 28.17
	ATOM		75.476 50.717 19.268 1.00 26.95
	ATOM		77.524 52.595 15.824 1.00 45.82
20	ATOM		78.604 52.024 16.008 1.00 41.65
30	ATOM		77.422 53.936 15.782 1.00 48.88
	ATOM	283 N PRO 193	76.176 54.701 15.577 1.00 47.51
	ATOM	284 CD PRO 193	78.560 54.836 15.999 1.00 47.34
	ATOM	285 CA PRO 193	77.879 56.162 16.319 1.00 46.04
25	ATOM	286 CB PRO 193	76.675 56.126 15.438 1.00 46.24
35	ATOM	287 CG PRO 193	79.475 54.377 17.137 1.00 49.60
	ATOM	288 C PRO 193	79.000 54.033 18.218 1.00 54.05
	ATOM	289 O PRO 193 290 N ASP 194	80.783 54.383 16.891 1.00 50.63
	ATOM		81.769 53.951 17.885 1.00 54.57
40	ATOM	291 CA ASP 194	
40	ATOM	292 CB ASP 194	83.164 53.965 17.272 1.00 59.28
	ATOM	293 CG ASP 194	83.309 52.952 16.170 1.00 66.39
	ATOM	294 OD1 ASP 194	83.057 53.311 14.998 1.00 72.95
	ATOM	295 OD2 ASP 194	83.640 51.787 16.486 1.00 69.00
	ATOM	296 C ASP 194	81.769 54.726 19.198 1.00 54.41
45	ATOM	297 O ASP 194	82.229 54.221 20.222 1.00 55.27
	ATOM	298 N ASP 195	81.268 55.956 19.168 1.00 57.20
	ATOM	299 CA ASP 195	81.206 56.775 20.371 1.00 59.68
	ATOM	300 CB ASP 195	81.017 58.261 20.006 1.00 62.99
	ATOM	301 CG ASP 195	79.747 58.526 19.187 1.00 71.67
50	ATOM	302 OD1 ASP 195	78.734 58.956 19.796 1.00 70.17

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ATOM
            303 OD2 ASP 195
                                79.782 58.311 17.951 1.00 75.23
                              80.092 56.289 21.306 1.00 58.39
    ATOM
            304 C ASP 195
            305 O ASP 195
                              80.032 56.676 22.474 1.00 59.81
    ATOM
                              79.245 55.399 20.794 1.00 54.47
            306 N ILE 196
     ATOM
            307 CA ILE 196
                               78.141 54.840 21.568 1.00 49.00
    ATOM
5
                               76.839 54.780 20.731 1.00 46.64
            308 CB ILE 196
    ATOM
            309 CG2 ILE 196
                               75.701 54.195 21.560 1.00 42.11
    ATOM
                               76.467 56.184 20.241 1.00 44.23
    ATOM
           310 CG1 ILE 196
                               75.214 56.238 19.373 1.00 48.45
    ATOM 311 CD1 ILE 196
                              78.497 53.436 22.068 1.00 46.22
            312 C ILE 196
10
    ATOM
                              78.912 52.570 21.298 1.00 42.07
    ATOM
            313 O ILE 196
                               78.357 53.228 23.370 1.00 45.62
           314 N GLY 197
    ATOM
                                78.658 51.930 23.941 1.00 51.49
            315 CA GLY 197
    ATOM
                               80.005 51.832 24.625 1.00 54.64
    ATOM
            316 C GLY 197
                               80.377 50.759 25.092 1.00 49.98
            317 O GLY 197
15
    ATOM
            318 N GLN 198
                               80.726 52.946 24.725 1.00 60.08
    ATOM
            319 CA GLN 198
                                82.039 52.939 25.366 1.00 61.01
    ATOM
            320 CB GLN 198
                                83.082 53.568 24.441 1.00 55.55
    ATOM
            321 C GLN 198
                               82.044 53.633 26.733 1.00 59.57
     ATOM
                               83.103 54.016 27.232 1.00 61.30
            322 O GLN 198
20
     ATOM
            323 N SER 199
                               80.875 53.738 27.362 1.00 57.27
     ATOM
                               80.758 54.397 28.665 1.00 50.61
            324 CA SER 199
     ATOM
            325 CB SER 199
                               80.276 55.842 28.478 1.00 53.70
     ATOM
            326 OG SER 199
                               81.010 56.508 27.463 1.00 61.92
     ATOM
            327 C SER 199
                              79.848 53.684 29.675 1.00 46.41
25
     ATOM
            328 O SER 199
                               78.798 54.210 30.060 1.00 41.16
     ATOM
                               80.222 52.466 30.096 1.00 42.08
     ATOM
            329 N PRO 200
                                81.349 51.648 29.605 1.00 38.31
     ATOM
            330 CD PRO 200
                                79.409 51.722 31.065 1.00 44.04
     ATOM
            331 CA PRO 200
                                79,941 50,297 30,925 1.00 36.06
30
     ATOM
            332 CB PRO 200
                                81.377 50.504 30.583 1.00 37.43
     ATOM
            333 CG PRO 200
     ATOM
            334 C PRO 200
                               79.615 52.270 32.485 1.00 50.91
     ATOM
            335 O PRO 200
                               80.629 51.980 33.123 1.00 55.65
                              78.663 53.060 32.975 1.00 55.81
     ATOM
            336 N ILE 201
     ATOM
            337 CA ILE 201
                               78.781 53.651 34.311 1.00 57.24
35
            338 CB ILE 201
                               78.861 55.192 34.250 1.00 58.40
     ATOM
            339 CG2 ILE 201
                               80.218 55.622 33.709 1.00 60.49
     ATOM
                               77.716 55.751 33.404 1.00 62.42
            340 CG1 ILE 201
     ATOM
            341 CD1 ILE 201
                               77.819 57.234 33.137 1.00 61.68
     ATOM
            342 C ILE 201
                              77.728 53.241 35.332 1.00 56.52
40
     ATOM
            343 O ILE 201
                              77.961 53.352 36.537 1.00 60.89
     ATOM
                               76.564 52.794 34.871 1.00 52.76
            344 N VAL 202
     ATOM
            345 CA VAL 202
                                75,522 52,366 35,802 1.00 47,37
     ATOM
                                74,117 52.377 35.153 1.00 38.14
            346 CB VAL 202
     ATOM
                                73.092 51.804 36.117 1.00 30.35
            347 CG1 VAL 202
45
     ATOM
                                73.730 53.798 34.763 1.00 26.69
            348 CG2 VAL 202
     ATOM
           349 C VAL 202
                               75.885 50.958 36.285 1.00 53.65
     ATOM
            350 O VAL 202
                               75.914 50.010 35.500 1.00 55.10
     ATOM
            351 N SER 203
                               76.226 50.839 37.561 1.00 59.85
     ATOM
                               76.614 49.556 38.132 1.00 64.58
           352 CA SER 203
50
     ATOM
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WO 99/26966

PCT/US98/25296

	ATOM ATOM	353 CB SER 203 354 OG SER 203	77.209 49.749 39.532 1.00 68.95 78.396 50.523 39.483 1.00 74.02
	ATOM	355 C SER 203	75.493 48.528 38.197 1.00 61.69
	ATOM	356 O SER 203	74.351 48.846 38.535 1.00 63.63
5	ATOM	357 N MET 204	75.848 47.295 37.859 1.00 57.37
_	<b>ATOM</b>	358 CA MET 204	74.932 46.162 37.885 1.00 57.54
	ATOM	359 CB MET 204	74.847 45.505 36.501 1.00 56.59
	ATOM	360 CG MET 204	74.012 46.270 35.489 1.00 44.08
	ATOM	361 SD MET 204	72.255 46.228 35.884 1.00 46.62
10	ATOM	362 CE MET 204	71.775 44.758 35.013 1.00 48.37
	<b>ATOM</b>	363 C MET 204	75.522 45.178 38.888 1.00 55.86
	<b>ATOM</b>	364 O MET 204	76.746 45.089 39.027 1.00 58.94
	ATOM	365 N PRO 205	74.671 44.432 39.607 1.00 55.36
	<b>ATOM</b>	366 CD PRO 205	73.203 44.570 39.625 1.00 57.73
15	<b>ATOM</b>	367 CA PRO 205	75.119 43.453 40.604 1.00 56.82
	<b>ATOM</b>	368 CB PRO 205	73.814 43.042 41.295 1.00 59.79
	<b>ATOM</b>	369 CG PRO 205	72.769 43.281 40.255 1.00 57.85
	ATOM	370 C PRO 205	75.902 42.239 40.083 1.00 57.25
	<b>ATOM</b>	371 O PRO 205	75.683 41.118 40.541 1.00 66.28
20	<b>ATOM</b>	372 N ASP 206	76.822 42.462 39.147 1.00 58.75
	ATOM	373 CA ASP 206	77.639 41.389 38.586 1.00 61.09
	ATOM	374 CB ASP 206	76.802 40.462 37.685 1.00 66.07
	ATOM	375 CG ASP 206	76.158 41.190 36.521 1.00 70.97
	ATOM	376 OD1 ASP 206	74.989 41.613 36.662 1.00 76.97
25	ATOM	377 OD2 ASP 206	76.813 41.322 35.465 1.00 61.12
	ATOM	378 C ASP 206	78.865 41.910 37.832 1.00 61.96
	ATOM	379 O ASP 206	79.406 41.230 36.957 1.00 65.14
	ATOM	380 N GLY 207	79.282 43.130 38.158 1.00 63.00
	ATOM	381 CA GLY 207	80.455 43.709 37.522 1.00 64.43
30	ATOM	382 C GLY 207	80.224 44.467 36.229 1.00 64.81
	ATOM	383 O GLY 207	80.649 45.619 36.110 1.00 68.76
	ATOM	384 N ASP 208	79.584 43.827 35.253 1.00 63.53
	ATOM	385 CA ASP 208	79.316 44.459 33.962 1.00 58.96
	ATOM	386 CB ASP 208	78.746 43.434 32.974 1.00 62.84
35	ATOM	387 CG ASP 208	79.743 42.336 32.633 1.00 64.73 79.575 41.200 33.121 1.00 66.65
	ATOM	388 OD1 ASP 208	• • • •
	ATOM	389 OD2 ASP 208	80.701 42.610 31.878 1.00 68.91 78.368 45.646 34.110 1.00 56.65
	ATOM	390 C ASP 208	77.182 45.473 34.392 1.00 55.79
40	ATOM	391 O ASP 208	78.911 46.852 33.953 1.00 54.66
40	ATOM	392 N LYS 209	78.132 48.081 34.082 1.00 53.92
	ATOM ATOM	393 CA LYS 209 394 CB LYS 209	79.034 49.236 34.515 1.00 49.71
	ATOM	394 CB L 13 209 395 C LYS 209	77.395 48.420 32.785 1.00 48.30
	ATOM	395 C LTS 209 396 O LYS 209	77.767 47.945 31.711 1.00 45.62
45	ATOM	390 O L13 209 397 N VAL 210	76.367 49.258 32.894 1.00 43.87
43		397 N VAL 210 398 CA VAL 210	75.539 49.662 31.757 1.00 41.25
	ATOM ATOM	398 CA VAL 210 399 CB VAL 210	74.020 49.624 32.125 1.00 32.99
	ATOM	400 CG1 VAL 210	73.153 50.029 30.937 1.00 31.44
	ATOM	400 CG1 VAL 210 401 CG2 VAL 210	73.626 48.239 32.604 1.00 27.57
50	ATOM	401 CG2 VAL 210 402 C VAL 210	75.868 51.061 31.234 1.00 43.30
50	AIOM	402 C 4AL 210	, 5.000 51.001 51.05 · 1.00 · 5.50

ATOM 404 N ASP 211 ATOM 405 CA ASP 211 ATOM 406 CB ASP 211 ATOM 406 CB ASP 211 ATOM 407 CG ASP 211 ATOM 408 ODI ASP 211 ATOM 409 OD2 ASP 211 ATOM 410 C ASP 211 ATOM 411 O ASP 211 ATOM 411 O ASP 211 ATOM 412 N LEU 212 ATOM 413 CA LEU 212 ATOM 414 CB LEU 212 ATOM 415 CG LEU 212 ATOM 416 CDI LEU 212 ATOM 417 CD2 LEU 212 ATOM 418 C LEU 212 ATOM 419 O LEU 212 ATOM 410 C ASP 211 ATOM 411 CB LEU 212 ATOM 415 CG LEU 212 ATOM 416 CDI LEU 212 ATOM 417 CD2 LEU 212 ATOM 418 C LEU 212 ATOM 420 N GLU 213 ATOM 421 CA GLU 213 ATOM 421 CA GLU 213 ATOM 422 CB GLU 213 ATOM 424 CD GLU 213 ATOM 426 OE2 GLU 213 ATOM 427 C GLU 213 ATOM 428 O GLU 213 ATOM 430 CA ALA 214 ATOM 431 CB ALA 214 ATOM 432 C ALA 214 ATOM 433 CD PHE 215 ATOM 434 N PHE 215 ATOM 437 CC PHE 215 ATOM 438 CD1 PHE 215 ATOM 437 CC PHE 215 ATOM 438 CD1 PHE 215 ATOM 437 CC PHE 215 ATOM 438 CD1 PHE 215 ATOM 439 CD2 PHE 215 ATOM 440 CE1 PHE 215			100 0 3/47 010	76.261 51.951 31.994 1.00 44.65
ATOM 405 CA ASP 211 75.906 52.498 29.240 1.00 40.62 ATOM 406 CB ASP 211 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.686 52.232 27.943 1.00 43.49 76.886 52.232 27.943 1.00 43.49 76.180 54.427 72.092 1.00 42.13 78.111 53.549 26.574 1.00 37.49 76.180 54.427 72.092 1.00 42.53 74.491 53.001 28.921 1.00 44.56 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 43.76 74.006 53.982 29.684 1.00 44.57 74.006 53.982 29.684 1.00 44.56 74.004 416 CDL LEU 212 72.473 55.785 30.359 1.00 40.49 72.473 55.785 30.359 1.00 40.49 72.217 56.6923 32.551 1.00 40.49 72.217 56.6923 32.551 1.00 40.49 72.217 56.6923 32.551 1.00 40.49 72.217 56.6923 32.551 1.00 40.49 72.325 54.886 28.049 1.00 44.77 71.254 54.540 27.548 1.00 42.77 71.254 54.540 27.548 1.00 4				, •
ATOM 406 CB ASP 211 76.686 52.232 27.943 1.00 43.49 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.77 770.14 53.499 27.161 1.00 40.42 770.16 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.42.56 770.14 53.499 27.161 1.00 40.42.56 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.14 53.499 27.161 1.00 40.45 770.49 170.14 55.499 27.161 1.00 40.45 770.49 170.14 55.499 27.161 1.00 40.45 770.49 170.14 55.499 27.161 1.00 40.45 770.14 50.14				
5 ATOM 407 CG ASP 211 ATOM 408 OD1 ASP 211 ATOM 409 OD2 ASP 211 ATOM 410 C ASP 211 ATOM 410 C ASP 211 ATOM 411 O ASP 211 ATOM 412 N LEU 212 ATOM 413 CA LEU 212 ATOM 415 CG LEU 212 ATOM 416 CD1 LEU 212 ATOM 416 CD1 LEU 212 ATOM 419 O LEU 212 ATOM 419 O LEU 212 ATOM 419 C LEU 212 ATOM 419 C LEU 212 ATOM 410 C ASP 211 ATOM 411 C ASP 211 ATOM 412 R S LEU 212 ATOM 414 CB LEU 212 ATOM 415 CG LEU 212 ATOM 416 CD1 LEU 212 ATOM 417 CD2 LEU 212 ATOM 418 C LEU 212 ATOM 419 O LEU 213 ATOM 420 N GLU 213 ATOM 421 CA GLU 213 ATOM 422 CB GLU 213 ATOM 424 CD GLU 213 ATOM 425 OEI GLU 213 ATOM 426 OE2 GLU 213 ATOM 427 C GLU 213 ATOM 428 O GLU 213 ATOM 429 N ALA 214 ATOM 430 CA ALA 214 ATOM 431 CB ALA 214 ATOM 431 CB ALA 214 ATOM 432 C ALA 214 ATOM 433 CA ALA 214 ATOM 434 N PHE 215 ATOM 437 CG PHE 215 ATOM 437 CG PHE 215 ATOM 440 CEI PHE 215 ATOM 441 CE2 PHE 215 ATOM 442 CP BER 216 ATOM 444 O PHE 215 ATOM 445 N SER 216 ATOM 447 CB SER 216 ATOM 448 OG SER 216 ATOM 449 C SER 216 ATOM 440 OSER 216 ATOM 445 N SER 216 ATOM 445 ON SER 216 AT				
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ATOM 430 CA ALA 214 73.753 52.605 24.424 1.00 39.52 ATOM 431 CB ALA 214 74.952 51.726 24.740 1.00 35.16 74.952 71.726 24.740 1.00 35.16 72.460 51.852 24.694 1.00 37.14 ATOM 433 O ALA 214 71.795 51.390 23.767 1.00 42.29 ATOM 434 N PHE 215 72.098 51.773 25.970 1.00 31.60 ATOM 435 CA PHE 215 70.883 51.102 26.404 1.00 31.67 ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80 ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60 40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 446 CA SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 449 C SER 216 68.644 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				73.775 53.814 25.245 1.00 39.28
ATOM 431 CB ALA 214 74.952 51.726 24.740 1.00 35.16 ATOM 432 C ALA 214 72.460 51.852 24.694 1.00 37.14 ATOM 433 O ALA 214 71.795 51.390 23.767 1.00 42.29 ATOM 434 N PHE 215 72.098 51.773 25.970 1.00 31.60 ATOM 435 CA PHE 215 70.883 51.102 26.404 1.00 31.67 ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80 ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61 ATOM 449 C SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37			430 CA ALA 214	
ATOM 432 C ALA 214 72.460 51.852 24.694 1.00 37.14 ATOM 433 O ALA 214 71.795 51.390 23.767 1.00 42.29 ATOM 434 N PHE 215 72.098 51.773 25.970 1.00 31.60 ATOM 435 CA PHE 215 70.883 51.102 26.404 1.00 31.67 ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80 ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60 ATOM 442 CZ PHE 215 67.200 50.591 29.163 1.00 21.35 ATOM 443 C PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 444 O PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.608 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.608 55.249 25.165 1.00 43.86 ATOM 449 C SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				74.952 51.726 24.740 1.00 35.16
ATOM 433 O ALA 214 71.795 51.390 23.767 1.00 42.29 ATOM 434 N PHE 215 72.098 51.773 25.970 1.00 31.60 ATOM 435 CA PHE 215 70.883 51.102 26.404 1.00 31.67 ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80  35 ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 443 C PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 35.75 ATOM 445 N SER 216 68.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.608 55.249 25.165 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37	30		432 C ALA 214	72.460 51.852 24.694 1.00 37.14
ATOM 435 CA PHE 215 70.883 51.102 26.404 1.00 31.67 ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80  35 ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 68.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.608 55.249 25.165 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37		ATOM	433 O ALA 214	71.795 51.390 23.767 1.00 42.29
ATOM 435 CA PHE 215 70.883 51.102 26.404 1.00 31.67 ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80  35 ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 68.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.608 55.249 25.165 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37		ATOM	434 N PHE 215	
ATOM 436 CB PHE 215 70.728 51.217 27.922 1.00 24.80  ATOM 437 CG PHE 215 69.512 50.522 28.458 1.00 21.78  ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64  ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53  ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63  ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35  ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75  ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84  ATOM 445 N SER 216 68.506 53.739 25.001 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86  ATOM 448 OG SER 216 68.668 55.249 25.165 1.00 43.86  ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76  ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50  ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37			435 CA PHE 215	70.883 51.102 26.404 1.00 31.67
ATOM 438 CD1 PHE 215 69.512 50.522 28.458 1.00 21.78 ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.668 55.249 25.165 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37			436 CB PHE 215	70.728 51.217 27.922 1.00 24.80
ATOM 438 CD1 PHE 215 69.553 49.171 28.771 1.00 24.64 ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 68.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.608 55.249 25.165 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37	35		437 CG PHE 215	69.512 50.522 28.458 1.00 21.78
ATOM 439 CD2 PHE 215 68.328 51.223 28.658 1.00 21.53 ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37			438 CD1 PHE 215	69.553 49.171 28.771 1.00 24.64
ATOM 440 CE1 PHE 215 68.429 48.528 29.277 1.00 27.63 ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35 ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				68.328 51.223 28.658 1.00 21.53
ATOM 441 CE2 PHE 215 67.200 50.591 29.163 1.00 21.60  40 ATOM 442 CZ PHE 215 67.249 49.242 29.472 1.00 21.35  ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75  ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84  ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09  ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61  45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86  ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66  ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76  ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50  ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37			440 CE1 PHE 215	68.429 48.528 29.277 1.00 27.63
40       ATOM       442 CZ PHE       215       67.249       49.242       29.472       1.00 21.35         ATOM       443 C PHE       215       69.675       51.706       25.694       1.00 35.75         ATOM       444 O PHE       215       68.838       50.975       25.161       1.00 34.84         ATOM       445 N SER       216       69.604       53.035       25.665       1.00 39.09         ATOM       446 CA SER       216       68.506       53.739       25.001       1.00 40.61         45       ATOM       447 CB SER       216       68.668       55.249       25.165       1.00 43.86         ATOM       448 OG SER       216       68.616       55.603       26.537       1.00 68.66         ATOM       449 C SER       216       68.444       53.380       23.518       1.00 40.76         ATOM       451 N GLU       217       69.611       53.332       22.878       1.00 38.37		ATOM		67.200 50.591 29.163 1.00 21.60
ATOM 443 C PHE 215 69.675 51.706 25.694 1.00 35.75 ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61 45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37	40			
ATOM 444 O PHE 215 68.838 50.975 25.161 1.00 34.84 ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61 45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37	. •			69.675 51.706 25.694 1.00 35.75
ATOM 445 N SER 216 69.604 53.035 25.665 1.00 39.09 ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61 45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				
ATOM 446 CA SER 216 68.506 53.739 25.001 1.00 40.61 45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				69.604 53.035 25.665 1.00 39.09
45 ATOM 447 CB SER 216 68.668 55.249 25.165 1.00 43.86 ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				
ATOM 448 OG SER 216 68.616 55.603 26.537 1.00 68.66 ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37	45			
ATOM 449 C SER 216 68.444 53.380 23.518 1.00 40.76 ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37	. 2			
ATOM 450 O SER 216 67.362 53.161 22.969 1.00 35.50 ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				
ATOM 451 N GLU 217 69.611 53.332 22.878 1.00 38.37				67.362 53.161 22.969 1.00 35.50
				69.611 53.332 22.878 1.00 38.37
	50	ATOM	452 CA GLU 217	

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453 CB GLU 217
                                71.164 53.049 20.997 1.00 39.67
    ATOM
                                71.701 54.461 20.880 1.00 46.65
            454 CG GLU 217
    ATOM
            455 CD GLU 217
                                70.881 55.315 19.925 1.00 53.25
    ATOM
                                70.920 55.056 18.702 1.00 57.12
            456 OE1 GLU 217
    ATOM
                                70.189 56.240 20.400 1.00 54.13
            457 OE2 GLU 217
    ATOM
                               69.135 51.598 21.209 1.00 38.48
    ATOM
            458 C GLU 217
    ATOM 459 O GLU 217
                               68.416 51.378 20.228 1.00 43.00
    ATOM 460 N PHE 218
                               69.426 50.677 22.120 1.00 35.49
            461 CA PHE 218
                               68.934 49.313 22.018 1.00 31.76
    ATOM
                               69.743 48.392 22.925 1.00 29.10
10
    ATOM
            462 CB PHE 218
                               71.169 48.260 22.510 1.00 26.25
    ATOM
            463 CG PHE 218
    ATOM
            464 CD1 PHE 218
                                72.176 48.177 23.459 1.00 24.59
                                71.510 48.233 21.163 1.00 23.53
    ATOM
            465 CD2 PHE 218
            466 CE1 PHE 218
                                73.504 48.072 23.073 1.00 27.68
     ATOM
                                72.832 48.128 20.765 1.00 25.37
    ATOM
            467 CE2 PHE 218
15
    ATOM
            468 CZ PHE 218
                               73.834 48.047 21.721 1.00 28.43
            469 C PHE 218
                               67.445 49.202 22.321 1.00 31.30
    ATOM
            470 O PHE 218
                               66.726 48.496 21.621 1.00 35.18
    ATOM
            471 N THR 219
                               66.967 49.915 23.333 1.00 30.54
    ATOM
            472 CA THR 219
                               65.552 49.853 23.675 1.00 33.53
20
    ATOM
            473 CB THR 219
                               65.269 50.467 25.057 1.00 36.07
     ATOM
            474 OG1 THR 219
                                65.903 51.746 25.157 1.00 42.99
     ATOM
                                65.797 49.562 26.145 1.00 34.32
    ATOM
            475 CG2 THR 219
            476 C THR 219
                               64.680 50.514 22.609 1.00 34.53
     ATOM
                               63.507 50.162 22.450 1.00 36.57
    ATOM
            477 O THR 219
25
                               65.267 51.457 21.873 1.00 38.13
     ATOM
            478 N LYS 220
                               64.563 52.158 20.806 1.00 41.42
     ATOM
            479 CA LYS 220
            480 CB LYS 220
                               65.452 53.257 20.208 1.00 41.62
     ATOM
            481 C LYS 220
                               64.140 51.182 19.716 1.00 41.80
     ATOM
            482 O LYS 220
                               63.032 51.274 19.192 1.00 43.29
30
    ATOM
            483 N ILE 221
                              65.018 50.234 19.393 1.00 36.93
    ATOM
                               64.726 49.250 18.355 1.00 37.33
            484 CA ILE 221
     ATOM
            485 CB ILE 221
                               65.965 48.932 17.482 1.00 33.71
     ATOM
     ATOM 486 CG2 ILE 221
                               66.491 50.202 16.826 1.00 41.26
     ATOM 487 CG1 ILE 221
                               67.042 48.235 18.309 1.00 30.36
35
     ATOM 488 CD1 ILE 221
                               68.178 47.687 17.472 1.00 26.28
     ATOM 489 C ILE 221
                              64.141 47.922 18.845 1.00 40.49
                              63.593 47.159 18.048 1.00 43.43
            490 O ILE 221
     ATOM
                              64.219 47.651 20.144 1.00 39.43
     ATOM
            491 N ILE 222
                               63.703 46.394 20.667 1.00 35.49
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     ATOM
            492 CA ILE 222
            493 CB ILE 222
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            494 CG2 ILE 222
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	ATOM	565 O ASP 231	53.279 36.196 13.366 1.00 25.15
	ATOM	566 N PHE 232	55.364 36.490 14.170 1.00 22.29
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	ATOM	571 CD2 PHE 232	57.980 32.830 15.181 1.00 19.35
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	ATOM	573 CE2 PHE 232	58.359 31.496 15.114 1.00 20.63
	ATOM	574 CZ PHE 232	58.561 30.893 13.873 1.00 26.10
	ATOM	575 C PHE 232	55.018 34.093 14.328 1.00 23.51
	ATOM	576 O PHE 232	54.541 33.189 13.637 1.00 22.39
25	ATOM	577 N ALA 233	54.837 34.182 15.644 1.00 24.55
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	ATOM	581 O ALA 233	52.063 32.051 15.744 1.00 25.71
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	ATOM	584 CB LYS 234	50.136 35.838 15.229 1.00 30.40
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	ATOM	586 CD LYS 234	49.151 35.947 17.569 1.00 53.64
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55	ATOM	588 NZ LYS 234	46.773 35.268 18.060 1.00 54.22
	ATOM	589 C LYS 234	50.388 33.686 13.978 1.00 30.35
	ATOM	590 O LYS 234	49.318 33.142 13.716 1.00 32.50
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	ATOM	594 CG LYS 235	51.997 35.181 10.631 1.00 20.88
	ATOM	595 CD LYS 235	52.982 35.836 9.688 1.00 26.50
	ATOM	596 CE LYS 235	52.512 37.227 9.310 1.00 31.33
15		596 CE LTS 235 597 NZ LYS 235	53.439 37.862 8.341 1.00 36.51
45	ATOM ATOM	597 NZ L1S 235	51.508 31.554 11.791 1.00 28.37
		598 C LYS 235 599 O LYS 235	51.491 30.948 10.721 1.00 29.62
	ATOM		51.700 30.943 12.954 1.00 33.22
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50	ATOM		52.911 29.101 14.043 1.00 26.25
50	ATOM	602 CB LEU 236	32.911 29.101 14.043 1.00 20.23

PCT/US98/25296

## WO 99/26966

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5	ATOM	607 O LEU 236	50.013 29.342 14.588 1.00 34.23
•	ATOM	608 N PRO 237	49.811 28.134 12.695 1.00 44.89
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	ATOM	613 C PRO 237	48.278 27.072 14.430 1.00 49.12
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	ATOM	615 N MET 238	49,104 26.126 14.860 1.00 45.79
	ATOM	616 CA MET 238	49.029 25.558 16.200 1.00 52.79
15	ATOM	617 CB MET 238	50.133 24.505 16.378 1.00 49.72
13	ATOM	618 CG MET 238	49.861 23.195 15.637 1.00 58.16
	ATOM	619 SD MET 238	51.342 22.205 15.284 1.00 60.11
	ATOM	620 CE MET 238	50.993 21.626 13.625 1.00 53.03
	ATOM	621 C MET 238	49.103 26.593 17.324 1.00 53.36
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	ATOM	626 CG PHE 239	51.307 30.763 18.734 1.00 31.32
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	ATOM	628 CD2 PHE 239	52.425 31.402 20.790 1.00 29.45
	ATOM	629 CE1 PHE 239 630 CE2 PHE 239	50.970 32.973 19.682 1.00 32.29
	ATOM		51.828 32.659 20.737 1.00 26.00
20	ATOM	631 CZ PHE 239	48.590 29.592 18.344 1.00 37.40
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	ATOM		47.958 30.166 17.321 1.00 36.32
	ATOM	634 N SER 240	46.745 30.959 17.529 1.00 39.00
	ATOM	635 CA SER 240	46.385 31.724 16.258 1.00 47.52
25	ATOM	636 CB SER 240	47.390 32.671 15.947 1.00 52.67
35	ATOM	637 OG SER 240	47.390 32.671 13.947 1.00 32.67 45.539 30.158 18.032 1.00 36.82
	ATOM	638 C SER 240	44.548 30.743 18.485 1.00 43.02
	ATOM	639 O SER 240	45.617 28.833 17.931 1.00 38.98
	ATOM	640 N GLU 241 641 CA GLU 241	44.554 27.954 18.408 1.00 40.35
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40	ATOM	642 CB GLU 241	44.788 26.521 17.926 1.00 49.38
	ATOM	643 CG GLU 241	44.541 26.287 16.452 1.00 65.25
	ATOM	644 CD GLU 241	44.873 24.856 16.002 1.00 70.72
	ATOM	645 OE1 GLU 241	44.806 23.923 16.845 1.00 73.36
	ATOM	646 OE2 GLU 241	45.211 24.679 14.805 1.00 68.60
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            687 O ASP 246
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             718 CD2 LEU 250
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                                65.522 31.521 24.125 1.00 33.44
                                64.564 30.527 24.735 1.00 38.03
            766 CD GLU 257
     ATOM
                                 63.977 29.705 24.000 1.00 45.59
            767 OE1 GLU 257
     ATOM
                                 64.385 30.577 25.965 1.00 45.75
            768 OE2 GLU 257
     ATOM
15
            769 C GLU 257
                               65.595 33.526 20.840 1.00 21.68
     ATOM
                               66.586 33.421 20.107 1.00 20.02
            770 O GLU 257
     ATOM
                               64.383 33.852 20.391 1.00 17.07
     ATOM
            771 N ILE 258
            772 CA ILE 258
                               64.135 34.090 18.973 1.00 17.01
     ATOM
            773 CB ILE 258
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     ATOM
            774 CG2 ILE 258
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     ATOM
                                61.952 32.831 18.885 1.00 16.69
            775 CG1 ILE 258
     ATOM
                                60.450 32.783 18.632 1.00 16.31
            776 CD1 ILE 258
     ATOM
                               64,911 35.324 18.501 1.00 17.65
     ATOM
            777 C ILE 258
                               65.605 35.263 17.484 1.00 22.58
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     ATOM
            778 O ILE 258
            779 N MET 259
                                64.865 36.410 19.274 1.00 20.17
     ATOM
                                 65.584 37.628 18.909 1.00 15.03
             780 CA MET 259
     ATOM
             781 CB MET 259
                                 65.234 38.771 19.856 1.00 20.12
     ATOM
             782 CG MET 259
                                 63.791 39.191 19.775 1.00 17.19
     ATOM
                                63.523 40.795 20.524 1.00 28.92
             783 SD MET 259
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     ATOM
                                63.718 40.406 22.261 1.00 19.58
             784 CE MET 259
     ATOM
                                67.090 37.402 18.884 1.00 18.84
     ATOM
             785 C MET 259
                                67.783 37.912 17.996 1.00 29.07
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             786 O MET 259
                               67.590 36.618 19.837 1.00 21.45
     ATOM
             787 N SER 260
                                69.019 36.319 19.906 1.00 18.71
             788 CA SER 260
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     ATOM
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            789 CB SER 260
     ATOM
                                69.128 36.421 22.329 1.00 25.42
             790 OG SER 260
     ATOM
                               69.430 35.469 18.709 1.00 17.83
             791 C SER 260
     ATOM
             792 O SER 260
                               70.497 35.673 18.131 1.00 22.97
     ATOM
                               68.572 34.522 18.331 1.00 21.66
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             793 N LEU 261
                                68.837 33.663 17.179 1.00 20.98
             794 CA LEU 261
     ATOM
                                67.739 32.608 17.053 1.00 22.66
     ATOM
             795 CB LEU 261
                                67.719 31.759 15.781 1.00 22.12
             796 CG LEU 261
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             797 CD1 LEU 261
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     ATOM
             798 CD2 LEU 261
                               68.873 34.527 15.920 1.00 22.95
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             799 C LEU 261
                               69.779 34.402 15.091 1.00 22.62
             800 O LEU 261
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             801 N ARG 262
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     ATOM
     ATOM
            802 CA ARG 262
                                 67.816 36.301 14.643 1.00 25.32
                                 66.525 37.115 14.677 1.00 21.95
     ATOM
            803 CB ARG 262
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             804 CG ARG 262
     ATOM
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             805 CD ARG 262
     ATOM
                                62.990 36.377 13.599 1.00 22.18
             806 NE ARG 262
     ATOM
            807 CZ ARG 262
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                                 61.429 38.075 13.752 1.00 20.81
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            808 NH1 ARG 262
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            809 NH2 ARG 262
     ATOM
            810 C ARG 262
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            811 O ARG 262
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                               69.608 37.579 15.676 1.00 26.36
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                                70.818 38.400 15.705 1.00 27.02
            813 CA ALA 263
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     ATOM
                                70.997 39.045 17.087 1.00 25.80
            814 CB ALA 263
    ATOM
            815 C ALA 263
                               72.026 37.514 15.368 1.00 25.21
     ATOM
    ATOM
            816 O ALA 263
                               72.825 37.844 14.492 1.00 31.14
            817 N ALA 264
                               72.109 36.358 16.027 1.00 25.62
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             818 CA ALA 264
                                73.203 35.408 15.828 1.00 23.85
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                                73.062 34.237 16.794 1.00 17.15
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            819 CB ALA 264
            820 C ALA 264
                               73.345 34.901 14.391 1.00 26.03
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                               74.460 34.773 13.886 1.00 25.66
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            821 O ALA 264
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            822 N VAL 265
                               72.234 34.615 13.723 1.00 25.22
                                72.327 34.128 12.350 1.00 28.38
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     ATOM
            823 CA VAL 265
     ATOM
            824 CB VAL 265
                                71.028 33.457 11.857 1.00 24.59
            825 CG1 VAL 265
                                70.707 32.264 12.719 1.00 25.53
     ATOM
            826 CG2 VAL 265
     ATOM
                                69.881 34.440 11.853 1.00 20.86
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            827 C VAL 265
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     ATOM
            828 O VAL 265
                               73.024 34.973 10.222 1.00 34.75
                               72.795 36.464 11.896 1.00 30.10
            829 N ARG 266
     ATOM
     ATOM
            830 CA ARG 266
                                73.211 37.602 11.089 1.00 30.69
             831 CB ARG 266
                                72.170 38.713 11.148 1.00 25.13
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             832 CG ARG 266
                                70.976 38.406 10.299 1.00 25.43
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     ATOM
            833 CD ARG 266
                                69.999 39.537 10.277 1.00 29.56
     ATOM
            834 NE ARG 266
                                69.032 39.340 9.205 1.00 31.59
            835 CZ ARG 266
                                67.814 39.861 9.197 1.00 31.18
    ATOM
                                 67.408 40.611 10.215 1.00 31.01
     ATOM
            836 NH1 ARG 266
                                 67.012 39.648 8.163 1.00 28.21
     ATOM
            837 NH2 ARG 266
            838 C ARG 266
                               74.568 38.111 11.544 1.00 34.28
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     ATOM
                               74.877 39.300 11.423 1.00 41.19
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     ATOM
                               75.362 37.207 12.108 1.00 30.80
            840 N TYR 267
     ATOM
                                76.694 37.544 12.573 1.00 33.84
     ATOM
            841 CA TYR 267
                                77.202 36.461 13.534 1.00 32.56
     ATOM
            842 CB TYR 267
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     ATOM
            843 CG TYR 267
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             844 CD1 TYR 267
                                79.131 37.465 14.835 1.00 32.60
                                80.491 37.593 15.106 1.00 34.90
            845 CE1 TYR 267
     ATOM
             846 CD2 TYR 267
                                79.615 35.801 13.184 1.00 32.84
     ATOM
                                80.972 35.920 13.446 1.00 34.70
     ATOM
             847 CE2 TYR 267
                                81.404 36.816 14.405 1.00 36.21
45
     ATOM
            848 CZ TYR 267
                                82.749 36.940 14.651 1.00 39.48
     ATOM
            849 OH TYR 267
                               77.615 37.649 11.360 1.00 37.82
     ATOM
            850 C TYR 267
            851 O TYR 267
                               77.648 36.749 10.517 1.00 39.45
     ATOM
                               78.319 38.769 11.239 1.00 44.62
             852 N ASP 268
     ATOM
                               79.248 38.963 10.133 1.00 45.56
            853 CA ASP 268
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     ATOM
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WO 99/26966

PCT/US98/25296

	ATOM	854 CB ASP 268 855 CG ASP 268	79.096 40.366 9.533 1.00 46.62 80.068 40.624 8.391 1.00 50.96
	ATOM ATOM	856 OD1 ASP 268	80.204 39.755 7.502 1.00 55.65
	ATOM	857 OD2 ASP 268	80.700 41.699 8.384 1.00 52.09
5	ATOM	858 C ASP 268	80.675 38.751 10.630 1.00 44.44
,	ATOM	859 O ASP 268	81.242 39.614 11.304 1.00 45.68
	ATOM	860 N PRO 269	81.281 37.600 10.296 1.00 45.94
	ATOM	861 CD PRO 269	80.739 36.503 9.476 1.00 43.72
	ATOM	862 CA PRO 269	82.651 37.309 10.730 1.00 46.63
10	ATOM	863 CB PRO 269	82.884 35.889 10.208 1.00 43.88
10	ATOM	864 CG PRO 269	81.983 35.797 9.018 1.00 44.66
	ATOM	865 C PRO 269	83.682 38.298 10.190 1.00 50.80
	ATOM	866 O PRO 269	84.681 38.578 10.854 1.00 48.56
	ATOM	867 N ALA 270	83,407 38.858 9.012 1.00 55.09
15	ATOM	868 CA ALA 270	84.306 39.820 8.374 1.00 55.68
	ATOM	869 CB ALA 270	83.799 40.168 6.974 1.00 53.64
	ATOM	870 C ALA 270	84,528 41.096 9.196 1.00 56.18
	ATOM	871 O ALA 270	85.577 41.729 9.082 1.00 61.07
	ATOM	872 N SER 271	83.543 41.479 10.006 1.00 51.38
20	ATOM	873 CA SER 271	83.661 42.678 10.836 1.00 45.90
	<b>ATOM</b>	874 CB SER 271	82.710 43.774 10.346 1.00 44.49
	ATOM	875 OG SER 271	81.360 43.358 10.404 1.00 45.26
	ATOM	876 C SER 271	83.409 42.395 12.317 1.00 46.61
	ATOM	877 O SER 271	83.431 43.309 13.143 1.00 48.31
25	ATOM	878 N ASP 272	83.172 41.126 12.642 1.00 46.73
	ATOM	879 CA ASP 272	82.920 40.689 14.013 1.00 42.49
	ATOM	880 CB ASP 272	84.200 40.807 14.849 1.00 42.12
	ATOM	881 CG ASP 272	84.103 40.072 16.169 1.00 50.30
	ATOM	882 OD1 ASP 272	83.417 39.028 16.218 1.00 45.10
30	ATOM	883 OD2 ASP 272	84.708 40.537 17.160 1.00 57.61 81.769 41.465 14.658 1.00 40.95
	ATOM	884 C ASP 272	81.885 41.975 15.779 1.00 42.93
	ATOM	885 O ASP 272 886 N THR 273	80.651 41.531 13.945 1.00 38.57
	ATOM	886 N THR 273 887 CA THR 273	79.473 42.239 14.425 1.00 40.99
35	ATOM ATOM	888 CB THR 273	79.262 43.574 13.656 1.00 40.76
33	ATOM	889 OG1 THR 273	79.240 43.318 12.248 1.00 42.61
	ATOM	890 CG2 THR 273	80.373 44.574 13.965 1.00 39.67
	ATOM	891 C THR 273	78.210 41.397 14.251 1.00 39.94
	ATOM	892 O THR 273	78.202 40.419 13.494 1.00 36.66
40	ATOM	893 N LEU 274	77.168 41.757 14.993 1.00 36.08
	ATOM	894 CA LEU 274	75.867 41.096 14.907 1.00 34.28
	ATOM	895 CB LEU 274	75.343 40.699 16.292 1.00 30.96
	ATOM	896 CG LEU 274	75.952 39.536 17.068 1.00 30.19
	ATOM	897 CD1 LEU 274	75.310 39.472 18.444 1.00 26.29
45	ATOM	898 CD2 LEU 274	75.744 38.237 16.309 1.00 27.43
	ATOM	899 C LEU 274	74.943 42.163 14.347 1.00 36.49
	ATOM	900 O LEU 274	75.152 43.354 14.596 1.00 40.27
	ATOM	901 N THR 275	73.923 41.758 13.606 1.00 36.42
	ATOM	902 CA THR 275	72.994 42.731 13.062 1.00 35.07
50	ATOM	903 CB THR 275	72.773 42.522 11.556 1.00 36.04

	A TO 1 (	004 OC1 TID 275	74.028 42.625 10.875 1.00 41.52
	ATOM	904 OG1 THR 275 905 CG2 THR 275	71.852 43.583 11.008 1.00 36.47
	ATOM		71.673 42.655 13.814 1.00 34.32
	ATOM		71.055 41.590 13.907 1.00 34.96
_	ATOM		71.292 43.767 14.432 1.00 31.79
5	ATOM	908 N LEU 276	70.044 43.840 15.173 1.00 29.47
	ATOM	909 CA LEU 276	70.181 44.766 16.389 1.00 25.29
	ATOM	910 CB LEU 276	71.328 44.501 17.383 1.00 29.01
	ATOM	911 CG LEU 276	
	ATOM	912 CD1 LEU 276	
10	ATOM	913 CD2 LEU 276	71.358 43.042 17.834 1.00 22.79
	ATOM	914 C LEU 276	68.966 44.350 14.228 1.00 31.69
	ATOM	915 O LEU 276	69.175 45.335 13.510 1.00 33.87
	ATOM	916 N SER 277	67.862 43.608 14.162 1.00 33.07
	ATOM	917 CA SER 277	66.721 43.935 13.315 1.00 30.61
15	ATOM	918 CB SER 277	65.949 45.111 13.909 1.00 22.87
	ATOM	919 OG SER 277	65.587 44.822 15.250 1.00 23.35
	ATOM	920 C SER 277	67.103 44.200 11.860 1.00 31.85
	ATOM	921 O SER 277	66.433 44.958 11.158 1.00 32.13
	ATOM	922 N GLY 278	68.188 43.566 11.421 1.00 32.29
20	ATOM	923 CA GLY 278	68.664 43.716 10.058 1.00 37.59
	ATOM	924 C GLY 278	69.063 45.122 9.639 1.00 43.26
	ATOM	925 O GLY 278	69.313 45.358 8.455 1.00 42.60
	ATOM	926 N GLU 279	69.177 46.038 10.599 1.00 43.42
	ATOM	927 CA GLU 279	69.532 47.420 10.291 1.00 44.55
25	ATOM	928 CB GLU 279	68.292 48.310 10.394 1.00 44.66
	ATOM	929 CG GLU 279	67.671 48.344 11.783 1.00 54.19
	ATOM	930 CD GLU 279	66.400 49.171 11.845 1.00 64.96
	ATOM	931 OE1 GLU 279	65.627 49.174 10.859 1.00 71.43
	ATOM	932 OE2 GLU 279	66.167 49.814 12.891 1.00 66.65
30	ATOM	933 C GLU 279	70.654 48.019 11.133 1.00 45.52
	ATOM	934 O GLU 279	71.207 49.057 10.772 1.00 51.83
	ATOM	935 N MET 280	71.007 47.373 12.242 1.00 44.66
	ATOM	936 CA MET 280	72.060 47.904 13.105 1.00 34.22
	ATOM	937 CB MET 280	71.470 48.382 14.433 1.00 32.38
35	ATOM	938 CG MET 280	72.479 49.058 15.345 1.00 37.87
	ATOM	939 SD MET 280	71.912 49.201 17.052 1.00 41.78
	ATOM	940 CE MET 280	70.650 50.495 16.911 1.00 37.01
	ATOM	941 C MET 280	73.183 46.920 13.386 1.00 35.70
	ATOM	942 O MET 280	72.976 45.900 14.044 1.00 36.99
40	ATOM	943 N ALA 281	74.366 47.221 12.867 1.00 34.80
	ATOM	944 CA ALA 281	75.535 46.377 13.091 1.00 35.11
	ATOM	945 CB ALA 281	76.529 46.527 11.955 1.00 31.27
	ATOM	946 C ALA 281	76.155 46.837 14.406 1.00 35.96
	ATOM	947 O ALA 281	76.478 48.015 14.570 1.00 39.10
45	ATOM	948 N VAL 282	76.285 45.916 15.353 1.00 36.46
	ATOM	949 CA VAL 282	76.839 46.246 16.655 1.00 36.05
	ATOM	950 CB VAL 282	75.783 46.090 17.783 1.00 35.60
	ATOM	951 CG1 VAL 282	74.633 47.069 17.568 1.00 38.73
	ATOM	952 CG2 VAL 282	75.262 44.660 17.844 1.00 33.27
50	ATOM	953 C VAL 282	78.062 45.408 16.996 1.00 37.70

	ATOM	954 O VAL 282	78.137 44.223 16.660 1.00 37.45
	ATOM	955 N ALA 283	79.032 46.047 17.637 1.00 39.21
	ATOM	956 CA ALA 283	80.254 45.375 18.048 1.00 43.73
	ATOM	957 CB ALA 283	81.433 46.352 18.047 1.00 42.04
5	ATOM	958 C ALA 283	80.060 44.752 19.435 1.00 43.28
,	ATOM	959 O ALA 283	79.179 45.157 20.203 1.00 45.77
	ATOM	960 N ARG 284	80.903 43.774 19.744 1.00 41.96
	ATOM	961 CA ARG 284	80.866 43.044 21.004 1.00 44.87
	ATOM	962 CB ARG 284	82.084 42.125 21.087 1.00 46.34
10	ATOM	963 CG ARG 284	81.930 40.947 22.017 1.00 51.85
10	ATOM	964 CD ARG 284	83.107 40.010 21.844 1.00 60.73
	ATOM	965 NE ARG 284	83.262 39.571 20.455 1.00 54.30
	ATOM	966 CZ ARG 284	83.221 38.300 20.074 1.00 53.66
	ATOM	967 NH1 ARG 284	83.032 37.343 20.973 1.00 49.99
15	ATOM	968 NH2 ARG 284	83.379 37.984 18.797 1.00 47.31
	ATOM	969 C ARG 284	80.803 43.945 22.237 1.00 44.85
	ATOM	970 O ARG 284	79.896 43.806 23.062 1.00 48.26
	ATOM	971 N GLU 285	81.750 44.873 22.349 1.00 41.60
	ATOM	972 CA GLU 285	81.802 45.787 23.484 1.00 41.17
20	ATOM	973 CB GLU 285	83.043 46.675 23.392 1.00 39.97
	ATOM	974 C GLU 285	80.538 46.640 23.603 1.00 40.08
	ATOM	975 O GLU 285	80.023 46.849 24.703 1.00 41.16
	ATOM	976 N GLN 286	80.017 47.088 22.463 1.00 38.49
	ATOM	977 CA GLN 286	78.818 47.926 22.425 1.00 36.25
25	ATOM	978 CB GLN 286	78.549 48.401 20.997 1.00 39.50
	ATOM	979 CG GLN 286	79.619 49.311 20.424 1.00 43.62
	ATOM	980 CD GLN 286	79.324 49.710 18.987 1.00 49.48
	ATOM	981 OE1 GLN 286	79.253 48.856 18.097 1.00 48.41
	ATOM	982 NE2 GLN 286	79.125 51.000 18.755 1.00 47.15
30	ATOM	983 C GLN 286	77.563 47.255 22.988 1.00 35.40
	ATOM	984 O GLN 286	76.903 47.806 23.871 1.00 31.24
	ATOM	985 N LEU 287	77.234 46.071 22.480 1.00 32.96
	ATOM	986 CA LEU 287	76.055 45.349 22.950 1.00 33.40
	ATOM	987 CB LEU 287	75.767 44.138 22.054 1.00 28.67
35	ATOM	988 CG LEU 287	74.466 43.375 22.342 1.00 26.66
	ATOM	989 CD1 LEU 287	73.263 44.305 22.244 1.00 19.41
	ATOM	990 CD2 LEU 287	74.325 42.221 21.368 1.00 24.84
	ATOM	991 C LEU 287	76.234 44.914 24.406 1.00 34.81
	ATOM	992 O LEU 287	75.265 44.857 25.175 1.00 33.92
40	ATOM	993 N LYS 288	77.476 44.621 24.781 1.00 35.38
	ATOM	994 CA LYS 288	77.814 44.204 26.140 1.00 36.12
	ATOM	995 CB LYS 288	79.296 43.839 26.210 1.00 37.13 79.762 43.280 27.533 1.00 44.61
	ATOM	996 CG LYS 288	
4.5	ATOM	997 CD LYS 288	81.256 43.018 27.494 1.00 54.07 81.757 42.435 28.801 1.00 60.87
45	ATOM	998 CE LYS 288	81.757 42.433 28.801 1.00 60.87
	ATOM	999 NZ LYS 288	77.510 45.345 27.109 1.00 36.90
	ATOM	1000 C LYS 288	76.684 45.206 28.013 1.00 40.68
	ATOM	1001 O LYS 288 1002 N ASN 289	78.129 46.495 26.863 1.00 35.94
50	ATOM	1002 N ASN 289 1003 CA ASN 289	77,947 47.680 27.695 1.00 36.12
50	ATOM	1003 CA ASIN 209	77.777 77.000 27.073 1.00 30.12

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ATOM 1004 CB ASN 289
                                78.982 48.738 27.332 1.00 31.78
    ATOM 1005 CG ASN 289
                                80.388 48.263 27.569 1.00 40.31
    ATOM 1006 OD1 ASN 289
                                80.627 47.422 28.440 1.00 43.12
    ATOM 1007 ND2 ASN 289
                                 81.326 48.758 26.775 1.00 35.36
                               76,553 48.277 27.590 1.00 36.98
    ATOM 1008 C ASN 289
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    ATOM 1009 O ASN 289
                               76.099 48.959 28.509 1.00 34.29
                               75.883 48.032 26.466 1.00 32.65
    ATOM 1010 N GLY 290
                                74.541 48.550 26.256 1.00 28.61
    ATOM 1011 CA GLY 290
                               73.497 48.001 27.210 1.00 26.54
    ATOM 1012 C GLY 290
    ATOM 1013 O GLY 290
                               72.362 48.480 27.234 1.00 31.06
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    ATOM 1014 N GLY 291
                               73.861 46.978 27.977 1.00 28.89
    ATOM 1015 CA GLY 291
                                72.929 46.413 28.937 1.00 25.24
                               72.872 44.900 28.997 1.00 28.12
    ATOM 1016 C GLY 291
    ATOM 1017 O GLY 291
                               72.335 44.345 29.955 1.00 31.16
                               73.406 44.223 27.985 1.00 29.51
    ATOM 1018 N LEU 292
15
                                73.361 42.766 27.969 1.00 32.79
    ATOM 1019 CA LEU 292
                                73.304 42.240 26.531 1.00 28.00
    ATOM 1020 CB LEU 292
                                71.948 42.355 25.827 1.00 23.68
    ATOM 1021 CG LEU 292
                                72.004 41.626 24.509 1.00 26.12
    ATOM 1022 CD1 LEU 292
                                 70.851 41.764 26.694 1.00 23.36
    ATOM 1023 CD2 LEU 292
20
                               74.484 42.085 28.742 1.00 32.33
     ATOM 1024 C LEU 292
                               74.312 40.967 29.232 1.00 32.22
     ATOM 1025 O LEU 292
    ATOM 1026 N GLY 293
ATOM 1027 CA GLY 293
                                75.627 42.750 28.846 1.00 30.31
                                76.751 42.176 29.561 1.00 28.82
                               77.238 40.894 28.913 1.00 29.87
     ATOM 1028 C GLY 293
25
                                77.432 40.843 27.698 1.00 35.43
     ATOM 1029 O GLY 293
                                77.392 39.848 29.714 1.00 31.88
     ATOM 1030 N VAL 294
                                77.866 38.561 29.217 1.00 35.77
     ATOM 1031 CA VAL 294
                                78.232 37.590 30.363 1.00 34.29
     ATOM 1032 CB VAL 294
                                 79.462 38.092 31.095 1.00 37.54
     ATOM 1033 CG1 VAL 294
30
                                 77.065 37.425 31.322 1.00 25.62
     ATOM 1034 CG2 VAL 294
                                76.882 37.879 28.274 1.00 35.89
     ATOM 1035 C VAL 294
                                77.263 36.960 27.541 1.00 37.99
     ATOM 1036 O VAL 294
     ATOM 1037 N VAL 295
                                75.619 38.304 28.305 1.00 34.41
                                74.616 37.728 27.413 1.00 32.98
     ATOM 1038 CA VAL 295
35
                                73.208 38.298 27.677 1.00 31.25
     ATOM 1039 CB VAL 295
                                 72.208 37.706 26.694 1.00 23.54
     ATOM 1040 CG1 VAL 295
                                 72.783 37.993 29.101 1.00 23.07
     ATOM 1041 CG2 VAL 295
                                75.057 38.062 25.993 1.00 33.92
     ATOM 1042 C VAL 295
                                74.932 37.238 25.090 1.00 36.95
     ATOM 1043 O VAL 295
                                75.625 39.253 25.820 1.00 31.27
     ATOM 1044 N SER 296
                                76.118 39.695 24.521 1.00 33.38
     ATOM 1045 CA SER 296
     ATOM 1046 CB SER 296
                                76.667 41.115 24.620 1.00 24.78
                                77.368 41.478 23.449 1.00 25.43
     ATOM 1047 OG SER 296
                               77.216 38.748 24.045 1.00 35.86
     ATOM 1048 C SER 296
45
                                77.220 38.324 22.886 1.00 39.60
     ATOM 1049 O SER 296
                                78.135 38.402 24.943 1.00 37.41
     ATOM 1050 N ASP 297
                                79.227 37.490 24.602 1.00 35.39
     ATOM 1051 CA ASP 297
     ATOM 1052 CB ASP 297
                                80.147 37.269 25.808 1.00 43.07
     ATOM 1053 CG ASP 297
                                80.839 38.540 26.266 1.00 45.07
50
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WO 99/26966

PCT/US98/25296

	ATOM	1054 OD1 ASP 297	81.175 39.398 25.419 1.00 48.02
	ATOM	1055 OD2 ASP 297	81.064 38.670 27.485 1.00 50.13
	ATOM	1056 C ASP 297	78.662 36.145 24.161 1.00 30.87
	ATOM	1057 O ASP 297	79.155 35.534 23.213 1.00 33.92
5	ATOM	1058 N ALA 298	77.625 35.698 24.861 1.00 28.96
	<b>ATOM</b>	1059 CA ALA 298	76.971 34.428 24.574 1.00 30.60
	ATOM	1060 CB ALA 298	75.889 34.157 25.610 1.00 27.56
	ATOM	1061 C ALA 298	76.377 34.408 23.163 1.00 33.04
	ATOM	1062 O ALA 298	76.538 33.426 22.426 1.00 32.48
10	ATOM	1063 N ILE 299	75.706 35.493 22.786 1.00 30.92
	ATOM	1064 CA ILE 299	75.091 35.588 21.468 1.00 24.71
	ATOM	1065 CB ILE 299	74.138 36.789 21.368 1.00 22.98
	ATOM	1066 CG2 ILE 299	73.430 36.786 20.018 1.00 21.90
	ATOM	1067 CG1 ILE 299	73.091 36.707 22.477 1.00 20.91
15	ATOM	1068 CD1 ILE 299	72.266 37.951 22.634 1.00 19.86
	ATOM	1069 C ILE 299	76.168 35.680 20.395 1.00 26.77
	ATOM	1070 O ILE 299	76.036 35.069 19.335 1.00 30.21
	ATOM	1071 N PHE 300	77.238 36.428 20.673 1.00 29.08
	ATOM	1072 CA PHE 300	78.345 36.562 19.726 1.00 28.06
20	ATOM	1073 CB PHE 300	79.386 37.565 20.235 1.00 29.06
	ATOM	1074 CG PHE 300	79.289 38.920 19.590 1.00 28.14
	ATOM	1075 CD1 PHE 300	78.449 39.896 20.113 1.00 27.20
	ATOM	1076 CD2 PHE 300	80.017 39.209 18.437 1.00 29.11
	ATOM	1077 CE1 PHE 300	78.332 41.139 19.499 1.00 28.18
25	ATOM	1078 CE2 PHE 300	79.908 40.450 17.815 1.00 29.07
	ATOM	1079 CZ PHE 300	79.064 41.416 18.348 1.00 22.61
	ATOM	1080 C PHE 300	78.991 35.201 19.485 1.00 29.00
	ATOM	1081 O PHE 300	79.278 34.833 18.344 1.00 30.35
	ATOM	1082 N GLU 301	79.183 34.442 20.560 1.00 31.81
30	ATOM	1083 CA GLU 301	79.767 33.111 20.470 1.00 34.96
	ATOM	1084 CB GLU 301	79.962 32.528 21.865 1.00 30.78
	ATOM	1085 C GLU 301	78.850 32.210 19.634 1.00 35.49
	ATOM	1086 O GLU 301	79.322 31.438 18.793 1.00 35.76
26	ATOM	1087 N LEU 302	77.543 32.313 19.869 1.00 32.14
35	ATOM	1088 CA LEU 302	76.559 31.522 19.132 1.00 25.56
	ATOM	1089 CB LEU 302	75.147 31.760 19.682 1.00 23.33 73.992 31.006 19.010 1.00 28.73
	ATOM	1090 CG LEU 302	
	ATOM	1091 CD1 LEU 302	74.093 29.509 19.270 1.00 23.93 72.667 31.551 19.514 1.00 21.32
40	ATOM	1092 CD2 LEU 302	
40		1093 C LEU 302	76.617 31.885 17.650 1.00 23.10
	ATOM ATOM	1094 O LEU 302	76.664 31.001 16.796 1.00 26.79 76.672 33.181 17.353 1.00 22.79
		1095 N GLY 303 1096 CA GLY 303	76.745 33.631 17.333 1.00 22.79
	ATOM		76.743 33.031 13.974 1.00 21.00
15	ATOM	1097 C GLY 303	77.889 32.619 14.125 1.00 29.18
45	ATOM	1098 O GLY 303	79.132 33.182 15.912 1.00 31.15
	ATOM ATOM	1099 N ALA 304 1100 CA ALA 304	80.375 32.703 15.313 1.00 35.44
	ATOM	1100 CA ALA 304 1101 CB ALA 304	81.562 32.995 16.235 1.00 33.44
	ATOM	1101 CB ALA 304 1102 C ALA 304	80.300 31.208 14.978 1.00 35.15
50	ATOM	1102 C ALA 304 1103 O ALA 304	80.705 30.785 13.891 1.00 37.13
<b>J</b> U	ATOM	1103 U ALA 304	00.703 30.703 13.071 1.00 37.13

	ATOM	1104 N SER 305	79.753 30.414 15.892 1.00 33.91
	ATOM	1105 CA SER 305	79.638 28.979 15.663 1.00 36.39
	ATOM	1106 CB SER 305	79,395 28.237 16.980 1.00 32.71
	ATOM	1107 OG SER 305	78.265 28.749 17.663 1.00 48.66
5	ATOM	1108 C SER 305	78.558 28.619 14.641 1.00 37.61
•	ATOM	1109 O SER 305	78.747 27.697 13.845 1.00 39.92
	ATOM	1110 N LEU 306	77.443 29.349 14.651 1.00 38.21
	ATOM	1111 CA LEU 306	76.350 29.092 13.714 1.00 35.65
	ATOM	1112 CB LEU 306	75.094 29.894 14.077 1.00 25.49
10	ATOM	1113 CG LEU 306	74.209 29.374 15.212 1.00 26.18
10	ATOM	1114 CD1 LEU 306	72.988 30.262 15.361 1.00 23.40
	ATOM	1115 CD2 LEU 306	73.777 27.952 14.921 1.00 23.57
	ATOM	1116 C LEU 306	76.723 29.356 12.258 1.00 38.05
	ATOM	1117 O LEU 306	76.092 28.809 11.353 1.00 37.22
15	ATOM	1117 O EEO 300 1118 N SER 307	77.743 30.185 12.030 1.00 40.41
15	ATOM	1119 CA SER 307	78.199 30.511 10.677 1.00 40.85
	ATOM	1119 CA SER 307	79.415 31.442 10.736 1.00 37.32
	ATOM	1121 OG SER 307	79.086 32.678 11.344 1.00 56.20
	ATOM	1121 OG SER 307	78.550 29.270 9.852 1.00 39.87
20	ATOM	1122 C SER 307	78.221 29.191 8.670 1.00 44.27
20	ATOM	1124 N ALA 308	79.207 28.305 10.487 1.00 39.29
	ATOM	1125 CA ALA 308	79.609 27.066 9.826 1.00 33.10
	ATOM	1126 CB ALA 308	80.607 26.310 10.696 1.00 33.37
	ATOM	1127 C ALA 308	78.403 26.177 9.502 1.00 34.07
25	ATOM	1127 C ALA 308	78.467 25.340 8.600 1.00 40.61
25	ATOM	1129 N PHE 309	77.305 26.368 10.230 1.00 31.85
	ATOM	1130 CA PHE 309	76.095 25.581 10.015 1.00 35.24
	ATOM	1131 CB PHE 309	75.149 25.698 11.219 1.00 33.69
	ATOM	1132 CG PHE 309	75.618 24.954 12.437 1.00 36.16
30	ATOM	1133 CD1 PHE 309	76.785 25.327 13.090 1.00 43.79
30	ATOM	1134 CD2 PHE 309	74.903 23.867 12.922 1.00 38.03
	ATOM	1135 CE1 PHE 309	77.237 24.627 14.210 1.00 41.12
	ATOM	1136 CE2 PHE 309	75.346 23.161 14.040 1.00 41.08
	ATOM	1137 CZ PHE 309	76.514 23.543 14.683 1.00 38.37
35	ATOM	1138 C PHE 309	75.361 25.934 8.720 1.00 36.31
33	ATOM	1139 O PHE 309	74.633 25.095 8.173 1.00 37.84
	ATOM	1140 N ASN 310	75.567 27.155 8.225 1.00 35.22
	ATOM	1141 CA ASN 310	74.933 27.625 6.988 1.00 43.66
	ATOM	1142 CB ASN 310	75.536 26.930 5.760 1.00 54.13
40	ATOM	1143 CG ASN 310	76.980 27.339 5.501 1.00 68.29
40	ATOM	1144 OD1 ASN 310	77.297 28.527 5.412 1.00 74.62
	ATOM	1145 ND2 ASN 310	77.859 26.348 5.352 1.00 68.85
	ATOM	1146 C ASN 310	73.430 27.385 7.013 1.00 38.37
	ATOM	1147 O ASN 310	72.882 26.735 6.123 1.00 36.70
45	ATOM	1148 N LEU 311	72.780 27.865 8.062 1.00 35.22
73	ATOM	1149 CA LEU 311	71.345 27.690 8.206 1.00 34.32
	ATOM	1150 CB LEU 311	70.895 28.054 9.630 1.00 30.19
	ATOM	1150 CB LEU 311	71.458 27.306 10.845 1.00 26.76
	ATOM	1151 CO LEU 311	70.792 27.847 12.104 1.00 21.37
50	ATOM	1152 CD1 LEU 311	71.217 25.813 10.722 1.00 22.95
50	A I OIVI	1175 CDZ DDC 511	, 1,51, 50,0 10,0 1,00 35,00

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70.601 28.561 7.206 1.00 34.64
    ATOM 1154 C LEU 311
                              71.087 29.625 6.820 1.00 37.70
    ATOM 1155 O LEU 311
                              69.444 28.091 6.752 1.00 29.40
    ATOM 1156 N ASP 312
    ATOM 1157 CA ASP 312
                               68.634 28.867 5.823 1.00 28.65
    ATOM 1158 CB ASP 312
                               68.302 28.061 4.545 1.00 24.79
                               67.459 26.804 4.804 1.00 21.47
    ATOM 1159 CG ASP 312
                                66.994 26.549 5.932 1.00 27.92
    ATOM 1160 OD1 ASP 312
                                67.250 26.057 3.832 1.00 27.53
    ATOM 1161 OD2 ASP 312
                              67.380 29.346 6.557 1.00 25.92
    ATOM 1162 C ASP 312
                              67.167 28.985 7.717 1.00 26.98
    ATOM 1163 O ASP 312
10
                              66.540 30.122 5.878 1.00 21.78
    ATOM 1164 N ASP 313
                               65.315 30.653 6.471 1.00 22.89
    ATOM 1165 CA ASP 313
                               64.517 31.458 5.439 1.00 29.19
    ATOM 1166 CB ASP 313
    ATOM 1167 CG ASP 313
                               65.216 32.739 5.025 1.00 36.82
                                65.985 33.285 5.845 1.00 41.51
    ATOM 1168 OD1 ASP 313
15
    ATOM 1169 OD2 ASP 313
                                64.997 33.203 3.883 1.00 44.19
                              64.421 29.587 7.085 1.00 25.09
    ATOM 1170 C ASP 313
                               63.778 29.829 8.110 1.00 27.60
    ATOM 1171 O ASP 313
                               64.363 28.420 6.449 1.00 20.90
    ATOM 1172 N THR 314
                                63.538 27.322 6.942 1.00 22.71
    ATOM 1173 CA THR 314
20
                               63.408 26.208 5.884 1.00 22.07
    ATOM 1174 CB THR 314
                                62.825 26.746 4.693 1.00 23.15
    ATOM 1175 OG1 THR 314
                                62.542 25.079 6.401 1.00 18.17
    ATOM 1176 CG2 THR 314
                               64.080 26.734 8.249 1.00 19.95
    ATOM 1177 C THR 314
                               63.326 26.477 9.182 1.00 22.40
    ATOM 1178 O THR 314
25
                               65.391 26.536 8.318 1.00 20.01
    ATOM 1179 N GLU 315
                                65.997 25.987 9.523 1.00 19.40
    ATOM 1180 CA GLU 315
                                67.454 25.626 9.254 1.00 11.72
    ATOM 1181 CB GLU 315
                                67.544 24.440 8.322 1.00 13.43
    ATOM 1182 CG GLU 315
    ATOM 1183 CD GLU 315
                                68.925 24.157 7.791 1.00 18.51
30
                                69.666 25.107 7.451 1.00 23.24
    ATOM 1184 OE1 GLU 315
    ATOM 1185 OE2 GLU 315
                                69.254 22.962 7.673 1.00 24.23
     ATOM 1186 C GLU 315
                               65.833 26.960 10.681 1.00 20.12
                               65.425 26.570 11.777 1.00 20.53
    ATOM 1187 O GLU 315
    ATOM 1188 N VAL 316
                               66.055 28.240 10.406 1.00 21.79
35
                                65.898 29.270 11.425 1.00 18.14
    ATOM 1189 CA VAL 316
                                66.346 30.659 10.898 1.00 18.97
     ATOM 1190 CB VAL 316
     ATOM 1191 CG1 VAL 316
                                66.040 31.741 11.929 1.00 19.08
     ATOM 1192 CG2 VAL 316
                                67.840 30.641 10.537 1.00 17.97
                               64,430 29.332 11.880 1.00 22.54
     ATOM 1193 C VAL 316
40
                               64.146 29.433 13.072 1.00 26.47
     ATOM 1194 O VAL 316
                               63.505 29.242 10.924 1.00 19.66
     ATOM 1195 N ALA 317
                                62.076 29.286 11.216 1.00 16.99
     ATOM 1196 CA ALA 317
                                61.279 29.329 9.926 1.00 17.79
     ATOM 1197 CB ALA 317
                               61.619 28.105 12.063 1.00 14.12
     ATOM 1198 C ALA 317
45
                               60.808 28.263 12.970 1.00 17.04
     ATOM 1199 O ALA 317
                               62.104 26.911 11.740 1.00 20.37
     ATOM 1200 N LEU 318
     ATOM 1201 CA LEU 318
                                61.725 25.714 12.485 1.00 21.12
                                62.131 24.448 11.718 1.00 21.80
     ATOM 1202 CB LEU 318
     ATOM 1203 CG LEU 318
                                61.364 24.265 10.398 1.00 18.11
50
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ATOM 1210 CB LEU 319 ATOM 1211 CG LEU 319 ATOM 1212 CD1 LEU 319 ATOM 1213 CD2 LEU 319 ATOM 1213 CD2 LEU 319 ATOM 1215 O LEU 319 ATOM 1215 O LEU 319 ATOM 1216 N GLN 320 ATOM 1217 CA GLN 320 ATOM 1218 CB GLN 320 ATOM 1219 CG GLN 320 ATOM 1219 CG GLN 320 ATOM 1220 CD GLN 320 ATOM 1221 OEI GLN 320 ATOM 1221 OEI GLN 320 ATOM 1222 NE2 GLN 320 ATOM 1224 OEI GLN 320 ATOM 1225 N ALA 321 ATOM 1226 CA ALA 321 ATOM 1229 C ALA 321 ATOM 1230 N VAL 322 ATOM 1231 CA VAL 322 ATOM 1231 CA VAL 322 ATOM 1233 CGI VAL 322 ATOM 1234 CG2 VAL 322 ATOM 1235 C VAL 322 ATOM 1236 C VAL 322 ATOM 1237 N LEU 323 ATOM 1240 CG LEU 323 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 323 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 323 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 323 ATOM 1242 CD2 LEU 323 ATOM 1242 CD2 LEU 323 ATOM 1244 CD2 LEU 323 ATOM 1245 N LEU 323 ATOM 1246 CA LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1250 CD2 LEU 324		ATOM	1204 CD1 LEU 318	61.946 23.125 9.594 1.00 16.79
ATOM 1206 C LEU 318 62.335 25.752 13.880 1.00 22.03 ATOM 1207 O LEU 318 61.688 25.373 14.858 1.00 21.35   ATOM 1208 N LEU 319 63.564 26.257 13.964 1.00 20.03   ATOM 1210 CB LEU 319 64.260 26.395 15.236 1.00 20.2   ATOM 1211 CG LEU 319 66.567 26.960 15.001 1.00 19.0   ATOM 1211 CD1 LEU 319 66.567 26.960 15.001 1.00 19.0   ATOM 1212 CD1 LEU 319 66.518 25.883 17.083 1.00 29.3   ATOM 1214 C LEU 319 66.518 25.883 17.083 1.00 29.3   ATOM 1215 O LEU 319 66.518 25.883 17.083 1.00 29.3   ATOM 1216 N GLN 320 68.012 27.326 15.699 1.00 20.5   ATOM 1216 N GLN 320 62.957 31.496 15.118 1.00 21.16   ATOM 1217 CA GLN 320 62.958 28.439 15.539 1.00 20.5   ATOM 1219 CG GLN 320 62.957 31.496 15.311 1.00 21.0   ATOM 1220 CD GLN 320 62.657 32.617 14.150 1.00 22.8   ATOM 1221 OE1 GLN 320 62.637 32.617 14.150 1.00 22.8   ATOM 1222 NE2 GLN 320 63.574 33.537 14.006 1.00 20.5   ATOM 1223 C GLN 320 60.368 28.976 17.844 1.00 23.3   ATOM 1224 O GLN 320 60.368 28.976 17.844 1.00 23.3   ATOM 1225 N ALA 321 60.251 27.886 15.876 1.00 22.7   ATOM 1227 CB ALA 321 59.010 27.187 16.201 1.00 19.8   ATOM 1230 N VAL 322 60.368 25.561 17.396 1.00 20.2   ATOM 1231 CA VAL 322 ATOM 1233 CG VAL 322 ATOM 1235 C VAL 322 ATOM 1235 C VAL 322 ATOM 1236 CO VAL 322 ATOM 1237 N LEU 323 ATOM 1238 CA LEU 323 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 323 ATOM 1242 CD2 LEU 323 ATOM 1244 C LEU 323 ATOM 1244 C LEU 323 ATOM 1244 C LEU 323 ATOM 1245 N LEU 323 ATOM 1246 CA LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1246 CA LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1246 CA LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG		ATOM	1205 CD2 LEU 318	
ATOM 1207 O LEU 318 5 ATOM 1208 N LEU 319 ATOM 1209 CA LEU 319 ATOM 1210 CB LEU 319 ATOM 1211 CG LEU 319 ATOM 1211 CG LEU 319 ATOM 1212 CD1 LEU 319 ATOM 1213 CD2 LEU 319 ATOM 1214 C LEU 319 ATOM 1215 O LEU 319 ATOM 1215 O LEU 319 ATOM 1216 N GLN 320 ATOM 1217 CA GLN 320 ATOM 1217 CA GLN 320 ATOM 1218 CB GLN 320 ATOM 1219 CG GLN 320 ATOM 1220 CD GLN 320 ATOM 1221 OE1 GLN 320 ATOM 1220 NE2 GLN 320 ATOM 1221 OE1 GLN 320 ATOM 1221 OE1 GLN 320 ATOM 1222 NE2 GLN 320 ATOM 1223 C GLN 320 ATOM 1224 O GLN 320 ATOM 1225 N ALA 321 ATOM 1226 CA ALA 321 ATOM 1227 CB ALA 321 ATOM 1231 CA VAL 322 ATOM 1231 CA VAL 322 ATOM 1232 CB VAL 322 ATOM 1234 CG2 VAL 322 ATOM 1235 C VAL 322 ATOM 1235 C VAL 322 ATOM 1236 O VAL 322 ATOM 1237 N LEU 323 ATOM 1237 N LEU 323 ATOM 1239 CB LEU 323 ATOM 1240 CG LEU 323 ATOM 1240 CG LEU 323 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 323 ATOM 1242 CD2 LEU 323 ATOM 1242 CD2 LEU 323 ATOM 1244 C LEU 323 ATOM 1246 CA LEU 323 ATOM 1247 CB LEU 323 ATOM 1247 CB LEU 323 ATOM 1248 CG LEU 323 ATOM 1249 CG LEU 323 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1246 CA LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1250 CD2 LEU 324				
5 ATOM 1208 N LEU 319 ATOM 1209 CA LEU 319 ATOM 1210 CB LEU 319 ATOM 1210 CB LEU 319 ATOM 1211 CG LEU 319 ATOM 1212 CD1 LEU 319 ATOM 1212 CD1 LEU 319 ATOM 1213 CD2 LEU 319 ATOM 1214 C LEU 319 ATOM 1215 O LEU 319 ATOM 1216 N GLN 320 ATOM 1217 CA GLN 320 ATOM 1219 CG GLN 320 ATOM 1220 CD GLN 320 ATOM 1221 CD1 GLN 320 ATOM 1221 CD1 GLN 320 ATOM 1219 CG GLN 320 ATOM 1221 OCD GLN 320 ATOM 1221 OCD GLN 320 ATOM 1221 OCD GLN 320 ATOM 1222 NE2 GLN 320 ATOM 1223 C GLN 320 ATOM 1225 N ALA 321 ATOM 1226 CA ALA 321 ATOM 1227 CB ALA 321 ATOM 1229 C ALA 321 ATOM 1230 N VAL 322 ATOM 1230 CB VAL 322 ATOM 1231 CA VAL 322 ATOM 1234 CG2 VAL 322 ATOM 1234 CG2 VAL 322 ATOM 1235 C VAL 322 ATOM 1236 O VAL 322 ATOM 1237 CB LEU 323 ATOM 1238 CA LEU 323 ATOM 1238 CA LEU 323 ATOM 1238 CA LEU 323 ATOM 1240 CD LEU 323 ATOM 1241 CD LEU 323 ATOM 1240 CD LEU 323 ATOM 1231 CA VAL 322 ATOM 1234 CG2 VAL 322 ATOM 1235 C VAL 322 ATOM 1236 O VAL 322 ATOM 1237 N LEU 323 ATOM 1240 CD LEU 323 ATOM 1241 CD LEU 323 ATOM 1240 CD LEU 323 ATOM 1241 CD LEU 323 ATOM 1240 CD LEU 323 ATOM 1241 CD LEU 323 ATOM 1240 CD LEU 323 ATOM 1241 CD LEU 324 ATOM 1240 CD LEU 323 ATOM 1241 CD LEU 324 ATOM 1240 CD LEU 324 ATOM 1245 N LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 C REU 324 ATOM 1249 CD LEU 324 ATOM 1240 CD LEU 324 ATOM 1240 CD LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 C REU 324 ATOM 1249 CD LEU 324 ATOM 1250 C		ATOM		61.688 25.373 14.858 1.00 21.35
ATOM 1209 CA LEU 319 ATOM 1210 CB LEU 319 ATOM 1211 CG LEU 319 ATOM 1212 CD1 LEU 319 ATOM 1212 CD1 LEU 319 ATOM 1213 CD2 LEU 319 ATOM 1214 C LEU 319 ATOM 1215 CD2 LEU 319 ATOM 1216 N GLN 320 ATOM 1216 N GLN 320 ATOM 1217 CA GLN 320 ATOM 1219 CG GLN 320 ATOM 1219 CG GLN 320 ATOM 1219 CG GLN 320 ATOM 1220 CD GLN 320 ATOM 1221 OE1 GLN 320 ATOM 1221 OE1 GLN 320 ATOM 1221 OE1 GLN 320 ATOM 1222 NE2 GLN 320 ATOM 1222 NE2 GLN 320 ATOM 1225 N ALA 321 ATOM 1225 N ALA 321 ATOM 1226 CA ALA 321 ATOM 1229 O ALA 321 ATOM 1229 O ALA 321 ATOM 1230 N VAL 322 ATOM 1231 CA VAL 322 ATOM 1233 CG1 VAL 322 ATOM 1233 CG1 VAL 322 ATOM 1234 CG2 VAL 322 ATOM 1235 C VAL 322 ATOM 1236 CA ALEU 323 ATOM 1237 N LEU 323 ATOM 1240 CG LEU 323 ATOM 1237 N LEU 323 ATOM 1240 CG LEU 323 ATOM 1237 N LEU 323 ATOM 1240 CG LEU 323 ATOM 1240 CG LEU 323 ATOM 1240 CG LEU 323 ATOM 1237 N LEU 323 ATOM 1236 CA LEU 323 ATOM 1240 CG LEU 324 ATOM 1240 CG LEU 323 ATOM 1240 CG LEU 324 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 324 ATOM 1240 CG LEU 323 ATOM 1241 CD1 LEU 324 ATOM 1245 N LEU 324 ATOM 1246 CA LEU 324 ATOM 1247 CB LEU 324 ATOM 1248 CG LEU 324 ATOM 1249 CD1 LEU 324 ATOM 1250 CD2 LEU 324 ATOM 12	5			
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ATOM 1241 CD1 LEU 323 64.719 29.532 19.528 1.00 14.3 ATOM 1242 CD2 LEU 323 64.816 27.750 21.320 1.00 21.3  40 ATOM 1243 C LEU 323 60.398 27.932 21.410 1.00 22.55 ATOM 1244 O LEU 323 60.185 27.986 22.615 1.00 25.21 ATOM 1245 N LEU 324 59.507 28.300 20.502 1.00 24.15 ATOM 1246 CA LEU 324 58.200 28.827 20.855 1.00 19.8 ATOM 1247 CB LEU 324 57.499 29.384 19.608 1.00 15.2 ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2 ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.3 ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.00				
ATOM 1242 CD2 LEU 323 64.816 27.750 21.320 1.00 21.3  40 ATOM 1243 C LEU 323 60.398 27.932 21.410 1.00 22.55  ATOM 1244 O LEU 323 60.185 27.986 22.615 1.00 25.21  ATOM 1245 N LEU 324 59.507 28.300 20.502 1.00 24.15  ATOM 1246 CA LEU 324 58.200 28.827 20.855 1.00 19.8  ATOM 1247 CB LEU 324 57.499 29.384 19.608 1.00 15.2  ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2  ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.9  ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.60		ATOM		
40       ATOM       1243       C       LEU       323       60.398       27.932       21.410       1.00       22.55         ATOM       1244       O       LEU       323       60.185       27.986       22.615       1.00       25.21         ATOM       1245       N       LEU       324       59.507       28.300       20.502       1.00       24.15         ATOM       1247       CB       LEU       324       58.200       28.827       20.855       1.00       19.8         45       ATOM       1248       CG       LEU       324       56.067       29.908       19.767       1.00       17.2         ATOM       1249       CD1       LEU       324       56.021       31.161       20.637       1.00       15.9         ATOM       1250       CD2       LEU       324       55.496       30.208       18.395       1.00       20.63		ATOM		
ATOM 1244 O LEU 323 60.185 27.986 22.615 1.00 25.21 ATOM 1245 N LEU 324 59.507 28.300 20.502 1.00 24.15 ATOM 1246 CA LEU 324 58.200 28.827 20.855 1.00 19.8 ATOM 1247 CB LEU 324 57.499 29.384 19.608 1.00 15.2 ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2 ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.5 ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.60				
ATOM 1245 N LEU 324 59.507 28.300 20.502 1.00 24.15 ATOM 1246 CA LEU 324 58.200 28.827 20.855 1.00 19.8 ATOM 1247 CB LEU 324 57.499 29.384 19.608 1.00 15.2 ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2 ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.5 ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.60	40	ATOM		
ATOM 1246 CA LEU 324 58.200 28.827 20.855 1.00 19.8 ATOM 1247 CB LEU 324 57.499 29.384 19.608 1.00 15.2 45 ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2 ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.5 ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.6				
ATOM 1247 CB LEU 324 57.499 29.384 19.608 1.00 15.2  45 ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2  ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.5  ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.6				
45 ATOM 1248 CG LEU 324 56.067 29.908 19.767 1.00 17.2 ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.5 ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.0				
ATOM 1249 CD1 LEU 324 56.021 31.161 20.637 1.00 15.9 ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.0				
ATOM 1250 CD2 LEU 324 55.496 30.208 18.395 1.00 20.0	45			
ATOM 1251 C I FII 324 57 311 27 795 21 536 1 00 19 83				
		ATOM		57.311 27.795 21.536 1.00 19.83
				56.767 28.064 22.609 1.00 24.47
50 ATOM 1253 N MET 325 57.197 26.603 20.956 1.00 25.02	50	ATOM	1253 N MET 325	57.197 26.603 20.956 1.00 25.02

	ATOM	1254 CA MET 325	56.339 25.563 21.522 1.00 26.72
	ATOM	1255 CB MET 325	55.823 24.644 20.410 1.00 30.03
	ATOM	1256 CG MET 325	55.129 25.358 19.241 1.00 25.09
	ATOM	1257 SD MET 325	53.714 26.409 19.672 1.00 27.29
5	ATOM	1258 CE MET 325	52.503 25.220 20.084 1.00 20.67
	ATOM	1259 C MET 325	56.995 24.736 22.635 1.00 28.94
	ATOM	1260 O MET 325	56.881 23.510 22.672 1.00 32.94
	ATOM	1261 N SER 326	57.642 25.418 23.569 1.00 29.36
	ATOM	1262 CA SER 326	58.311 24.759 24.680 1.00 31.62
10	ATOM	1263 CB SER 326	59.554 25.559 25.064 1.00 38.13
	ATOM	1264 OG SER 326	60.277 24.949 26.119 1.00 48.99
	ATOM	1265 C SER 326	57.361 24.653 25.871 1.00 33.69
	ATOM	1266 O SER 326	56.620 25.594 26.166 1.00 33.66
	ATOM	1267 N THR 327	57.356 23.499 26.536 1.00 38.27
15	ATOM	1268 CA THR 327	56.497 23.306 27.701 1.00 38.98
	ATOM	1269 CB THR 327	55.875 21.896 27.730 1.00 33.30
	ATOM	1270 OG1 THR 327	56.908 20.911 27.627 1.00 44.01
	ATOM	1271 CG2 THR 327	54.888 21.722 26.587 1.00 38.09
_	ATOM	1272 C THR 327	57.239 23.570 29.018 1.00 42.88
20	ATOM	1273 O THR 327	56.702 23.325 30.099 1.00 43.36
	ATOM	1274 N ASP 328	58.462 24.091 28.924 1.00 45.92
	ATOM	1275 CA ASP 328	59.268 24.410 30.104 1.00 49.59
	ATOM	1276 CB ASP 328	60.760 24.411 29.760 1.00 59.87
0.5	ATOM	1277 CG ASP 328	61.273 23.040 29.387 1.00 75.73
25	ATOM	1278 OD1 ASP 328	62.008 22.939 28.382 1.00 85.81
	ATOM	1279 OD2 ASP 328	60.946 22.063 30.098 1.00 85.56
	ATOM	1280 C ASP 328	58.873 25.767 30.673 1.00 48.50
	ATOM	1281 O ASP 328	59.725 26.609 30.961 1.00 57.50
20	ATOM	1282 N ARG 329 1283 CA ARG 329	57.569 25.980 30.805 1.00 49.62 57.032 27.222 31.340 1.00 50.52
30	ATOM ATOM	1283 CA ARG 329 1284 CB ARG 329	56.400 28.080 30.230 1.00 53.57
	ATOM	1285 CG ARG 329	57.376 28.828 29.324 1.00 51.09
	ATOM	1286 CD ARG 329	57.897 27.951 28.204 1.00 49.73
	ATOM	1287 NE ARG 329	58.692 28.699 27.233 1.00 47.44
35	ATOM	1288 CZ ARG 329	60.005 28.569 27.080 1.00 54.28
33	ATOM	1289 NH1 ARG 329	60.688 27.722 27.839 1.00 58.35
	ATOM	1290 NH2 ARG 329	60.631 29.256 26.136 1.00 51.92
	ATOM	1291 C ARG 329	55.970 26.870 32.375 1.00 51.90
	ATOM	1292 O ARG 329	55.378 25.790 32.324 1.00 50.77
40	ATOM	1293 N SER 330	55.728 27.784 33.303 1.00 50.56
	ATOM	1294 CA SER 330	54.744 27.564 34.349 1.00 50.67
	ATOM	1295 CB SER 330	55.271 28.108 35.678 1.00 46.64
	ATOM	1296 C SER 330	53.404 28.213 34.004 1.00 47.63
	ATOM	1297 O SER 330	53.371 29.309 33.440 1.00 48.02
45	ATOM	1298 N GLY 331	52.314 27.496 34.277 1.00 44.44
	ATOM	1299 CA GLY 331	50.977 28.023 34.044 1.00 38.77
	ATOM	1300 C GLY 331	50.236 27.710 32.756 1.00 41.74
	ATOM	1301 O GLY 331	49.147 28.246 32.537 1.00 49.57
	ATOM	1302 N LEU 332	50.783 26.841 31.912 1.00 39.75
50	ATOM	1303 CA LEU 332	50.123 26.502 30.651 1.00 37.55

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51.107 25.829 29.694 1.00 32.36
    ATOM 1304 CB LEU 332
    ATOM 1305 CG LEU 332
                               52.268 26.659 29.153 1.00 34.40
    ATOM 1306 CD1 LEU 332
                                53.207 25.749 28.379 1.00 30.22
                                51.742 27.786 28.277 1.00 23.33
    ATOM 1307 CD2 LEU 332
                               48.921 25.589 30.834 1.00 36.73
    ATOM 1308 C LEU 332
                               48.987 24.608 31.577 1.00 39.29
    ATOM 1309 O LEU 332
    ATOM 1310 N LEU 333
                               47.822 25.925 30.168 1.00 36.07
                               46.615 25.107 30.215 1.00 39.58
    ATOM 1311 CA LEU 333
    ATOM 1312 CB LEU 333
                               45.384 25.906 29.754 1.00 41.08
    ATOM 1313 CG LEU 333
                                44.601 26.883 30.644 1.00 47.59
10
    ATOM 1314 CD1 LEU 333
                               44.268 26.213 31.961 1.00 45.65
                                45.366 28.171 30.874 1.00 47.42
    ATOM 1315 CD2 LEU 333
                               46.791 23.911 29.278 1.00 40.00
    ATOM 1316 C LEU 333
                               46.690 22.754 29.689 1.00 44.77
    ATOM 1317 O LEU 333
    ATOM 1318 N CYA 334
                               47.102 24.213 28.022 1.00 37.70
                                47.265 23.209 26.968 1.00 36.04
    ATOM 1319 CA CYA 334
                                46.815 23.808 25.635 1.00 40.64
    ATOM 1320 CB CYA 334
                                45.280 24.738 25.758 1.00 44.31
    ATOM 1321 SG CYA 334
    ATOM 1322 AS CYA 334
                                43.972 22.946 25.380 1.00 76.30
20
    ATOM 1323 C CYA 334
                               48.668 22.617 26.815 1.00 34.91
    ATOM 1324 O CYA 334
                               49.237 22.615 25.722 1.00 37.63
    ATOM 1325 N VAL 335
                               49.189 22.056 27.903 1.00 35.43
                               50.518 21.452 27.909 1.00 34.27
    ATOM 1326 CA VAL 335
                               50.861 20.868 29.298 1.00 34.21
    ATOM 1327 CB VAL 335
                                52.261 20.258 29.292 1.00 33.66
    ATOM 1328 CG1 VAL 335
25
                                50.755 21.945 30.362 1.00 31.77
    ATOM 1329 CG2 VAL 335
                               50.662 20.349 26.865 1.00 37.14
    ATOM 1330 C VAL 335
    ATOM 1331 O VAL 335
                               51.639 20.320 26.114 1.00 37.59
    ATOM 1332 N ASP 336
                               49.683 19.451 26.813 1.00 39.99
    ATOM 1333 CA ASP 336
                               49.705 18.339 25.866 1.00 41.64
30
    ATOM 1334 CB ASP 336
                               48.532 17.392 26.146 1.00 54.27
                                48.596 16.118 25.322 1.00 67.42
    ATOM 1335 CG ASP 336
                               47.915 16.049 24.274 1.00 70.98
    ATOM 1336 OD1 ASP 336
    ATOM 1337 OD2 ASP 336
                              49.337 15.191 25.717 1.00 76.88
                              49.702 18.762 24.393 1.00 38.31
    ATOM 1338 C ASP 336
35
    ATOM 1339 O ASP 336
                               50.469 18.229 23.586 1.00 37.46
    ATOM 1340 N LYS 337
                               48.853 19.729 24.052 1.00 30.23
    ATOM 1341 CA LYS 337
                               48.740 20.211 22.676 1.00 29.21
                               47.561 21.189 22.559 1.00 30.53
    ATOM 1342 CB LYS 337
    ATOM 1343 CG LYS 337
                                47.012 21.360 21.162 1.00 51.63
40
                               45.636 21.997 21.186 1.00 59.57
    ATOM 1344 CD LYS 337
    ATOM 1345 CE LYS 337
                               45.066 22.115 19.774 1.00 66.05
                               43.673 22.693 19.776 1.00 67.20
    ATOM 1346 NZ LYS 337
    ATOM 1347 C LYS 337
                               50.054 20.873 22.249 1.00 28.33
                               50.581 20.594 21.170 1.00 26.08
    ATOM 1348 O LYS 337
45
    ATOM 1349 N ILE 338
                              50.609 21.696 23.141 1.00 26.74
                               51.873 22.390 22.902 1.00 25.42
    ATOM 1350 CA ILE 338
    ATOM 1351 CB ILE 338
                               52.177 23.379 24.052 1.00 23.57
    ATOM 1352 CG2 ILE 338
                               53.559 23.991 23.874 1.00 22.59
                               51.105 24.471 24.096 1.00 23.57
50
    ATOM 1353 CG1 ILE 338
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51.157 25.362 25.333 1.00 24.30
    ATOM 1354 CD1 ILE 338
    ATOM 1355 C ILE 338
                              53.018 21.382 22.768 1.00 29.20
    ATOM 1356 O ILE 338
                              53.905 21.537 21.916 1.00 31.59
                               52,977 20.340 23.595 1.00 34.82
    ATOM 1357 N GLU 339
                                53.980 19.277 23.597 1.00 34.23
    ATOM 1358 CA GLU 339
                                53.639 18.256 24.681 1.00 40.38
    ATOM 1359 CB GLU 339
                                54.785 17.354 25.072 1.00 54.98
    ATOM 1360 CG GLU 339
                                55.644 17.964 26.178 1.00 71.26
    ATOM 1361 CD GLU 339
    ATOM 1362 OE1 GLU 339
                                56.766 18.444 25.858 1.00 77.82
                                55.170 17.985 27.349 1.00 65.14
    ATOM 1363 OE2 GLU 339
10
    ATOM 1364 C GLU 339
                               53.972 18.582 22.231 1.00 34.42
                               55.018 18.431 21.590 1.00 29.41
    ATOM 1365 O GLU 339
                               52.778 18.189 21.786 1.00 34.13
    ATOM 1366 N LYS 340
                               52.592 17.513 20.502 1.00 32.05
    ATOM 1367 CA LYS 340
    ATOM 1368 CB LYS 340
                               51.121 17.105 20.325 1.00 34.59
15
    ATOM 1369 C LYS 340
                               53.064 18.390 19.337 1.00 32.56
    ATOM 1370 O LYS 340
                               53.762 17.913 18.441 1.00 32.93
                               52.725 19.677 19.374 1.00 31.42
    ATOM 1371 N SER 341
                              53.134 20.621 18.334 1.00 27.79
    ATOM 1372 CA SER 341
    ATOM 1373 CB SER 341
                               52.559 22.009 18.601 1.00 27.85
20
                               51.149 21.966 18.579 1.00 47.20
    ATOM 1374 OG SER 341
    ATOM 1375 C SER 341
                               54.647 20.713 18.240 1.00 26.01
                               55.205 20.706 17.139 1.00 27.10
    ATOM 1376 O SER 341
                               55.318 20.794 19.389 1.00 24.25
    ATOM 1377 N GLN 342
                                56.771 20.875 19.392 1.00 27.16
    ATOM 1378 CA GLN 342
25
                                57,309 21,089 20,799 1.00 25.60
    ATOM 1379 CB GLN 342
    ATOM 1380 CG GLN 342
                                58.768 21.466 20.777 1.00 27.99
    ATOM 1381 CD GLN 342
                                59.407 21.429 22.133 1.00 29.58
    ATOM 1382 OE1 GLN 342
                                60.123 22.356 22.513 1.00 31.18
                                59.184 20.345 22.868 1.00 29.17
    ATOM 1383 NE2 GLN 342
30
    ATOM 1384 C GLN 342
                               57.377 19.609 18.786 1.00 28.45
                               58.378 19.675 18.062 1.00 29.79
    ATOM 1385 O GLN 342
    ATOM 1386 N GLU 343
                               56.777 18.458 19.078 1.00 26.58
    ATOM 1387 CA GLU 343
                                57.251 17.190 18.525 1.00 30.07
    ATOM 1388 CB GLU 343
                                56.462 16.016 19.114 1.00 40.79
35
     ATOM 1389 CG GLU 343
                                56.812 15.700 20.568 1.00 61.22
    ATOM 1390 CD GLU 343
                                55.951 14.594 21.166 1.00 71.76
                                55.472 13.719 20.405 1.00 76.73
    ATOM 1391 OE1 GLU 343
                                55.758 14.601 22.403 1.00 74.09
     ATOM 1392 OE2 GLU 343
                               57.097 17.225 17.001 1.00 25.87
    ATOM 1393 C GLU 343
40
     ATOM 1394 O GLU 343
                               58.008 16.842 16.260 1.00 27.26
     ATOM 1395 N ALA 344
                               55.947 17.727 16.550 1.00 23.70
     ATOM 1396 CA ALA 344
                                55.647 17.853 15.124 1.00 22.16
                                54.275 18.489 14.927 1.00 21.18
     ATOM 1397 CB ALA 344
                               56.729 18.694 14.454 1.00 21.24
     ATOM 1398 C ALA 344
45
     ATOM 1399 O ALA 344
                               57.303 18.284 13.438 1.00 26.47
                               57.048 19.840 15.055 1.00 22.48
     ATOM 1400 N TYR 345
     ATOM 1401 CA TYR 345
                                58.073 20.738 14.531 1.00 21.41
     ATOM 1402 CB TYR 345
                                58.085 22.059 15.304 1.00 20.10
     ATOM 1403 CG TYR 345
                                57.023 23.015 14.830 1.00 15.87
50
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ATOM 1404 CD1 TYR 345
                                56.004 23.434 15.682 1.00 10.54
                                54.983 24.259 15.225 1.00 17.09
    ATOM 1405 CE1 TYR 345
                                57.003 23.448 13.505 1.00 16.86
    ATOM 1406 CD2 TYR 345
                                55.991 24.269 13.036 1.00 16.84
     ATOM 1407 CE2 TYR 345
                                54.984 24.668 13.896 1.00 17.97
    ATOM 1408 CZ TYR 345
                                53.963 25.455 13.406 1.00 27.11
    ATOM 1409 OH TYR 345
                               59.465 20.120 14.548 1.00 24.43
    ATOM 1410 C TYR 345
                               60.238 20.291 13.597 1.00 24.69
    ATOM 1411 O TYR 345
                               59.777 19.401 15.621 1.00 26.75
    ATOM 1412 N LEU 346
                                61.074 18.746 15.767 1.00 25.06
    ATOM 1413 CA LEU 346
10
                                61.207 18.108 17.150 1.00 24.59
    ATOM 1414 CB LEU 346
                                61.637 19.076 18.252 1.00 26.46
    ATOM 1415 CG LEU 346
                                61.387 18.468 19.610 1.00 26.46
     ATOM 1416 CD1 LEU 346
                                63.101 19.437 18.076 1.00 21.78
     ATOM 1417 CD2 LEU 346
                               61.322 17.713 14.683 1.00 23.24
     ATOM 1418 C LEU 346
15
                               62.416 17.645 14.127 1.00 27.54
     ATOM 1419 O LEU 346
                               60.314 16.900 14.395 1.00 25.75
     ATOM 1420 N LEU 347
                                60.437 15.881 13.356 1.00 25.41
     ATOM 1421 CA LEU 347
                                59.208 14.970 13.330 1.00 23.78
     ATOM 1422 CB LEU 347
                                59.302 13.713 14.190 1.00 31.85
     ATOM 1423 CG LEU 347
20
                                58.004 12.928 14.089 1.00 39.88
     ATOM 1424 CD1 LEU 347
                                 60.483 12.864 13.738 1.00 27.65
     ATOM 1425 CD2 LEU 347
                               60.611 16.535 11.998 1.00 23.22
     ATOM 1426 C LEU 347
                               61.468 16.133 11.211 1.00 28.58
     ATOM 1427 O LEU 347
     ATOM 1428 N ALA 348
                                59.784 17.542 11.731 1.00 26.40
25
                                59.840 18.273 10.474 1.00 23.85
     ATOM 1429 CA ALA 348
                                58.732 19.324 10.433 1.00 25.27
     ATOM 1430 CB ALA 348
                                61.210 18.924 10.337 1.00 23.69
     ATOM 1431 C ALA 348
                                61.847 18.835 9.288 1.00 29.11
     ATOM 1432 O ALA 348
                               61.678 19.506 11.438 1.00 24.71
     ATOM 1433 N PHE 349
30
                                62.973 20.181 11.493 1.00 20.48
     ATOM 1434 CA PHE 349
                                63.164 20.772 12.900 1.00 17.84
     ATOM 1435 CB PHE 349
                                64.334 21.721 13.031 1.00 14.90
     ATOM 1436 CG PHE 349
                                65.109 22.069 11.933 1.00 17.58
     ATOM 1437 CD1 PHE 349
                                 64.651 22.269 14.271 1.00 24.77
     ATOM 1438 CD2 PHE 349
35
                                 66.185 22.944 12.063 1.00 20.26
     ATOM 1439 CE1 PHE 349
                                 65.727 23.147 14.413 1.00 23.83
     ATOM 1440 CE2 PHE 349
                                66.494 23.486 13.299 1.00 20.36
     ATOM 1441 CZ PHE 349
                                64.084 19.181 11.159 1.00 23.43
     ATOM 1442 C PHE 349
                                64.916 19.427 10.278 1.00 24.35
     ATOM 1443 O PHE 349
40
                                64.057 18.028 11.820 1.00 25.79
     ATOM 1444 N GLU 350
                                 65.060 16.991 11.606 1.00 26.75
     ATOM 1445 CA GLU 350
                                 64.813 15.822 12.567 1.00 29.56
     ATOM 1446 CB GLU 350
                                 65.774 14.661 12.391 1.00 39.94
     ATOM 1447 CG GLU 350
                                 65.574 13.549 13.407 1.00 45.06
     ATOM 1448 CD GLU 350
45
                                 64.413 13.192 13.715 1.00 49.26
     ATOM 1449 OE1 GLU 350
                                 66.593 13.017 13.887 1.00 56.67
     ATOM 1450 OE2 GLU 350
                                65.051 16.494 10.162 1.00 26.95
     ATOM 1451 C GLU 350
                                66.096 16.398 9.513 1.00 28.77
     ATOM 1452 O GLU 350
                               63.858 16.219 9.652 1.00 22.56
     ATOM 1453 N HIS 351
50
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63.699 15.728 8.294 1.00 22.20
    ATOM 1454 CA HIS 351
                               62.263 15.265 8.083 1.00 22.47
    ATOM 1455 CB HIS 351
                               61.881 14.106 8.947 1.00 23.61
    ATOM 1456 CG HIS 351
                               62.633 13.300 9.739 1.00 27.65
    ATOM 1457 CD2 HIS 351
                                60.585 13.653 9.069 1.00 26.13
    ATOM 1458 ND1 HIS 351
                               60.548 12.629 9.898 1.00 22.87
    ATOM 1459 CE1 HIS 351
                               61.779 12.393 10.319 1.00 27.53
    ATOM 1460 NE2 HIS 351
                              64.135 16.764 7.259 1.00 21.76
    ATOM 1461 C HIS 351
                              64.708 16.419 6.226 1.00 27.02
    ATOM 1462 O HIS 351
                               63.909 18.041 7.555 1.00 18.26
    ATOM 1463 N TYR 352
10
                                64.327 19.101 6.649 1.00 16.94
    ATOM 1464 CA TYR 352
                                63.749 20.455 7.066 1.00 19.07
    ATOM 1465 CB TYR 352
                                64.107 21.534 6.081 1.00 21.11
    ATOM 1466 CG TYR 352
    ATOM 1467 CD1 TYR 352
                                63.518 21.564 4.819 1.00 21.33
                                63.921 22.482 3.859 1.00 21.06
    ATOM 1468 CE1 TYR 352
15
    ATOM 1469 CD2 TYR 352
                                65.105 22.462 6.367 1.00 22.07
                                65.515 23.388 5.412 1.00 25.40
    ATOM 1470 CE2 TYR 352
                                64.921 23.384 4.161 1.00 21.90
     ATOM 1471 CZ TYR 352
                                65.334 24.268 3.197 1.00 23.57
     ATOM 1472 OH TYR 352
                               65.853 19.156 6.657 1.00 18.49
     ATOM 1473 C TYR 352
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     ATOM 1474 O TYR 352
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     ATOM 1475 N VAL 353
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     ATOM 1476 CA VAL 353
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     ATOM 1479 CG2 VAL 353
                               68.452 17.829 7.146 1.00 25.07
     ATOM 1480 C VAL 353
     ATOM 1481 O VAL 353
ATOM 1482 N ASN 354
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                               67.768 16.690 7.221 1.00 24.59
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                                67.223 14.331 6.751 1.00 26.05
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     ATOM 1486 OD1 ASN 354
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                               68.143 15.813 4.981 1.00 30.50
     ATOM 1488 C ASN 354
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                                66.926 16.901 3.157 1.00 26.02
     ATOM 1491 CA HIS 355
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     ATOM 1492 CB HIS 355
                                65.367 18.217 1.638 1.00 37.91
     ATOM 1493 CG HIS 355
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     ATOM 1495 ND1 HIS 355
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     ATOM 1497 NE2 HIS 355
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                               68.009 17.851 2.652 1.00 24.29
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     ATOM 1502 CB ARG 356
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     ATOM 1503 CG ARG 356
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    ATOM 1506 CZ ARG 356
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                                70.814 24.776 2.667 1.00 29.08
    ATOM 1507 NH1 ARG 356
    ATOM 1508 NH2 ARG 356
                                71.136 25.493 4.816 1.00 33.61
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                               71.140 17.937 3.519 1.00 30.56
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    ATOM 1512 CA LYS 357
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    ATOM 1514 CG LYS 357
                               71.674 14.676 0.550 1.00 54.23
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    ATOM 1516 CE LYS 357
    ATOM 1517 NZ LYS 357
                               69.288 13.974 0.162 1.00 65.88
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                               74.522 18.248 2.728 1.00 40.70
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                               74.863 19.581 5.163 1.00 33.59
    ATOM 1521 CA HIS 358
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    ATOM 1522 CB HIS 358
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    ATOM 1523 CG HIS 358
                               73.593 21.200 6.666 1.00 29.74
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    ATOM 1524 CD2 HIS 358
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    ATOM 1525 ND1 HIS 358
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    ATOM 1526 CE1 HIS 358
    ATOM 1527 NE2 HIS 358
                               71.747 22.373 6.838 1.00 23.32
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                              76.121 18.720 5.180 1.00 37.98
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                              76.087 17.581 5.654 1.00 41.07
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    ATOM 1536 CA ILE 360
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    ATOM 1538 CG2 ILE 360
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    ATOM 1539 CG1 ILE 360
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                              78.462 17.222 9.239 1.00 47.23
    ATOM 1541 C ILE 360
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                               80.869 16.705 9.729 1.00 50.93
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     ATOM 1545 CA PRO 361
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     ATOM 1546 CB PRO 361
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     ATOM 1547 CG PRO 361
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     ATOM 1548 C PRO 361
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    ATOM 1553 CG HIS 362
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    ATOM 1565 CD2 PHE 363
                                72.596 20.076 10.575 1.00 27.51
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     ATOM 1567 CE2 PHE 363
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    ATOM 1569 C PHE 363
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     ATOM 1580 CZ2 TRP 364
     ATOM 1581 CZ3 TRP 364
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                                76.131 19.681 15.945 1.00 27.18
                                77.341 18.802 16.149 1.00 23.71
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     ATOM 1596 CD LYS 366
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     ATOM 1597 CE LYS 366
                                79.624 18.329 15.231 1.00 35.88
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                                74.383 19.497 21.545 1.00 50.01
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           1622 CE MET 369
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     ATOM 1645 CG2 THR 372
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     ATOM 1653 OD2 ASP 373
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     ATOM 1702 CB CYA 380
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     ATOM 1703 SG CYA 380
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WO 99/26966

WO 99/26966

# PCT/US98/25296

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	ATOM	1706 O CYA 380 70.337 37.033 32.270 1.00 45.73
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•	ATOM	1709 CB HIS 381 68.377 37.687 28.565 1.00 25.76
	ATOM	1710 CG HIS 381 67.596 38.932 28.285 1.00 20.30
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20	ATOM	1724 CB SER 383 69.233 36.297 35.719 1.00 42.58
	ATOM	1725 OG SER 383 68.734 35.010 36.049 1.00 61.85
	ATOM	1726 C SER 383 68.638 38.697 35.311 1.00 36.49
	ATOM	1727 O SER 383 68.443 39.480 36.243 1.00 43.81
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	ATOM	1731 CG ARG 384 71.590 41.679 32.731 1.00 29.20
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	ATOM	1737 C ARG 384 68.777 41.431 33.916 1.00 39.45
35	ATOM	1738 O ARG 384 68.913 42.537 34.444 1.00 44.47
55	ATOM	1739 N PHE 385 67.673 41.077 33.270 1.00 36.42
	ATOM	1740 CA PHE 385 66.568 42.007 33.099 1.00 34.68
	ATOM	1741 CB PHE 385 65.444 41.393 32.262 1.00 30.21
	ATOM	1742 CG PHE 385 64.263 42.304 32.081 1.00 29.48
40		1743 CD1 PHE 385 64.289 43.313 31.127 1.00 29.70
-70	ATOM	1744 CD2 PHE 385 63.130 42.161 32.873 1.00 28.04
	ATOM	1745 CE1 PHE 385 63.203 44.169 30.966 1.00 33.50
	ATOM	1746 CE2 PHE 385 62.040 43.012 32.718 1.00 31.35
	ATOM	1747 CZ PHE 385 62.077 44.017 31.763 1.00 32.08
45	ATOM	1748 C PHE 385 66.040 42.412 34.468 1.00 35.76
	ATOM	1749 O PHE 385 65.761 43.590 34.693 1.00 40.58
	ATOM	1750 N LEU 386 65.906 41.441 35.373 1.00 37.55
	ATOM	1751 CA LEU 386 65.429 41.706 36.735 1.00 41.01
	ATOM	1752 CB LEU 386 65.394 40.413 37.563 1.00 42.30
50	ATOM	1753 CG LEU 386 64.240 39.434 37.317 1.00 43.34
20	A I OIVI	1733 60 100 300 01.210 37.131 37.521 2.00 10131

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64,559 38,066 37,912 1.00 43.50
    ATOM 1754 CD1 LEU 386
                                62.946 39.992 37.899 1.00 44.01
    ATOM 1755 CD2 LEU 386
    ATOM 1756 C LEU 386
                               66.342 42.735 37.405 1.00 40.08
                               65.875 43.632 38.112 1.00 42.08
    ATOM 1757 O LEU 386
                              67.643 42.613 37.153 1.00 34.86
    ATOM 1758 N HIS 387
                               68.631 43.537 37.700 1.00 39.09
    ATOM 1759 CA HIS 387
                               70.046 43.034 37.421 1.00 39.99
    ATOM 1760 CB HIS 387
                               70.402 41.791 38.172 1.00 56.37
    ATOM 1761 CG HIS 387
                               71.384 40.881 37.974 1.00 60.11
    ATOM 1762 CD2 HIS 387
    ATOM 1763 ND1 HIS 387
                               69,711 41.370 39.290 1.00 60.40
10
                               70.252 40.255 39.746 1.00 61.89
    ATOM 1764 CE1 HIS 387
                               71.269 39.937 38.966 1.00 63.96
    ATOM 1765 NE2 HIS 387
                              68.446 44.928 37.101 1.00 41.00
    ATOM 1766 C HIS 387
                              68.492 45.927 37.817 1.00 46.99
    ATOM 1767 O HIS 387
                               68.213 44.982 35.792 1.00 39.15
    ATOM 1768 N MET 388
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    ATOM 1769 CA MET 388
                                68.011 46.243 35.088 1.00 35.32
    ATOM 1770 CB MET 388
                                67.676 45.992 33.612 1.00 35.12
                                68.810 45.442 32.753 1.00 37.24
    ATOM 1771 CG MET 388
    ATOM 1772 SD MET 388
                                68.259 45.150 31.051 1.00 41.75
                               69.274 43.748 30.573 1.00 35.23
    ATOM 1773 CE MET 388
20
                               66.880 47.048 35.733 1.00 36.52
    ATOM 1774 C MET 388
                               66.994 48.265 35.888 1.00 43.39
    ATOM 1775 O MET 388
                               65.792 46.371 36.103 1.00 38.05
    ATOM 1776 N LYS 389
                               64.637 47.025 36.729 1.00 42.88
    ATOM 1777 CA LYS 389
                               63.481 46.035 36.866 1.00 47.83
25
    ATOM 1778 CB LYS 389
                                62.835 45.627 35.560 1.00 52.36
    ATOM 1779 CG LYS 389
    ATOM 1780 CD LYS 389
                                62.040 44.340 35.731 1.00 61.84
                                60.978 44.451 36.814 1.00 69.04
    ATOM 1781 CE LYS 389
                                60.254 43.162 36.987 1.00 70.00
    ATOM 1782 NZ LYS 389
                               64.983 47.587 38.107 1.00 43.99
    ATOM 1783 C LYS 389
30
                               64.455 48.621 38.525 1.00 44.22
    ATOM 1784 O LYS 389
                               65.851 46.878 38.816 1.00 45.50
    ATOM 1785 N VAL 390
                                66.290 47.286 40.142 1.00 47.76
    ATOM 1786 CA VAL 390
                                67.152 46.186 40.804 1.00 46.30
    ATOM 1787 CB VAL 390
    ATOM 1788 CG1 VAL 390
                                67.796 46.706 42.079 1.00 49.20
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    ATOM 1789 CG2 VAL 390
                                66.305 44.962 41.097 1.00 42.69
                               67.109 48.571 40.070 1.00 47.25
    ATOM 1790 C VAL 390
                               66.811 49.540 40.760 1.00 48.67
    ATOM 1791 O VAL 390
    ATOM 1792 N GLU 391
                               68.115 48.580 39.199 1.00 44.11
                                69.009 49.721 39.047 1.00 45.79
     ATOM 1793 CA GLU 391
40
     ATOM 1794 CB GLU 391
                                70.266 49.311 38.273 1.00 45.78
    ATOM 1795 CG GLU 391
                                70.998 48.091 38.830 1.00 57.29
     ATOM 1796 CD GLU 391
                                71.479 48.268 40.261 1.00 61.20
     ATOM 1797 OE1 GLU 391
                                71.845 49.400 40.646 1.00 57.29
     ATOM 1798 OE2 GLU 391
                                71.496 47.263 41.001 1.00 63.69
45
                               68.410 50.959 38.391 1.00 49.16
     ATOM 1799 C GLU 391
     ATOM 1800 O GLU 391
                               68.463 52.055 38.956 1.00 58.82
     ATOM 1801 N CYA 392
                               67.802 50.782 37.224 1.00 49.75
                                67.255 51.908 36.475 1.00 45.56
     ATOM 1802 CA CYA 392
                                67.667 51.768 35.016 1.00 44.82
     ATOM 1803 CB CYA 392
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WO 99/26966

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69.443 51.771 34.913 1.00 50.78
    ATOM 1804 SG CYA 392
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     ATOM 1805 AS CYA 392
                               65.771 52.200 36.601 1.00 44.35
     ATOM 1806 C CYA 392
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    ATOM 1807 O CYA 392
                               65.378 53.469 36.365 1.00 45.52
     ATOM 1808 N PRO 393
                                66.275 54.603 36.075 1.00 37.38
     ATOM 1809 CD PRO 393
                                63.982 53.916 36.444 1.00 45.41
     ATOM 1810 CA PRO 393
     ATOM 1811 CB PRO 393
                                64.105 55.438 36.376 1.00 43.33
                                65.329 55.644 35.542 1.00 39.89
    ATOM 1812 CG PRO 393
    ATOM 1813 C PRO 393
                               63.108 53.376 35.318 1.00 44.89
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                               63.556 53.239 34.175 1.00 45.60
     ATOM 1814 O PRO 393
     ATOM 1815 N THR 394
                               61.843 53.135 35.647 1.00 47.52
    ATOM 1816 CA THR 394
                                60.853 52.603 34.713 1.00 53.06
    ATOM 1817 CB THR 394
                                59.459 52.583 35.371 1.00 61.06
                                59.609 52.470 36.794 1.00 72.44
     ATOM 1818 OG1 THR 394
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    ATOM 1819 CG2 THR 394
                                58.640 51.401 34.860 1.00 61.05
    ATOM 1820 C THR 394
                               60.767 53.373 33.392 1.00 49.98
    ATOM 1821 O THR 394
                               60.507 52.786 32.339 1.00 51.06
    ATOM 1822 N GLU 395
                               61.024 54.676 33.452 1.00 48.55
    ATOM 1823 CA GLU 395
                                60.970 55.548 32.282 1.00 44.21
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     ATOM 1824 CB GLU 395
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     ATOM 1825 C GLU 395
                               61.899 55.134 31.134 1.00 43.46
     ATOM 1826 O GLU 395
                               61.684 55.527 29.988 1.00 44.17
    ATOM 1827 N LEU 396
                               62.934 54.359 31.449 1.00 41.05
                                63.898 53.899 30.448 1.00 39.55
     ATOM 1828 CA LEU 396
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     ATOM 1829 CB LEU 396
                                65.270 53.708 31.106 1.00 35.03
     ATOM 1830 CG LEU 396
                                66.296 54.834 30.945 1.00 40.06
     ATOM 1831 CD1 LEU 396
                                65.638 56.200 31.055 1.00 39.06
                                67.398 54.669 31.978 1.00 32.78
     ATOM 1832 CD2 LEU 396
                               63.468 52.602 29.757 1.00 38.50
    ATOM 1833 C LEU 396
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    ATOM 1834 O LEU 396
                               64.106 52.150 28.804 1.00 34.72
                               62.364 52.028 30.225 1.00 38.76
     ATOM 1835 N PHE 397
     ATOM 1836 CA PHE 397
                                61.860 50.774 29.683 1.00 36.57
     ATOM 1837 CB PHE 397
                                61.610 49.775 30.819 1.00 33.96
                                62.842 49.421 31.607 1.00 36.95
    ATOM 1838 CG PHE 397
35
     ATOM 1839 CD1 PHE 397
                                63.331 50.280 32.587 1.00 34.61
                                63.523 48.234 31.362 1.00 37.14
     ATOM 1840 CD2 PHE 397
                                64.481 49.964 33.310 1.00 31.57
     ATOM 1841 CE1 PHE 397
                                64.675 47.908 32.082 1.00 37.85
    ATOM 1842 CE2 PHE 397
                               65.153 48.776 33.056 1.00 33.08
40
     ATOM 1843 CZ PHE 397
     ATOM 1844 C PHE 397
                               60.584 50.921 28.858 1.00 35.65
                               59.519 51.249 29.399 1.00 35.75
     ATOM 1845 O PHE 397
     ATOM 1846 N PRO 398
                               60.672 50.685 27.536 1.00 35.78
     ATOM 1847 CD PRO 398
                                61.891 50.367 26.767 1.00 32.81
                                59.503 50.786 26.658 1.00 33.94
     ATOM 1848 CA PRO 398
45
    ATOM 1849 CB PRO 398
                                60.041 50.297 25.315 1.00 33.91
                                61.488 50.707 25.356 1.00 33.09
     ATOM 1850 CG PRO 398
                               58.434 49.840 27.210 1.00 34.98
     ATOM 1851 C PRO 398
    ATOM 1852 O PRO 398
                               58.753 48.729 27.654 1.00 35.76
                               57.163 50.267 27.219 1.00 37.67
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     ATOM 1853 N PRO 399
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ATOM 1854 CD PRO 399
                                56.661 51.578 26.776 1.00 38.02
     ATOM 1855 CA PRO 399
                                56.070 49.433 27.733 1.00 36.86
     ATOM 1856 CB PRO 399
                                54.803 50.183 27.291 1.00 34.14
                                55.282 51.240 26.310 1.00 37.00
     ATOM 1857 CG PRO 399
     ATOM 1858 C PRO 399
                               56.085 47.970 27.273 1.00 37.06
                               55.967 47.063 28.099 1.00 37.07
     ATOM 1859 O PRO 399
                               56.299 47.738 25.980 1.00 35.13
     ATOM 1860 N LEU 400
                                56.327 46.374 25.445 1.00 35.86
     ATOM 1861 CA LEU 400
                                56.314 46.385 23.914 1.00 31.49
     ATOM 1862 CB LEU 400
                                56.181 45.017 23.227 1.00 30.73
     ATOM 1863 CG LEU 400
10
                                54.901 44.330 23.674 1.00 21.35
     ATOM 1864 CD1 LEU 400
                                56.197 45.183 21.720 1.00 25.42
     ATOM 1865 CD2 LEU 400
                               57.542 45.597 25.958 1.00 36.51
    ATOM 1866 C LEU 400
     ATOM 1867 O LEU 400
                               57.458 44.392 26.219 1.00 37.47
     ATOM 1868 N PHE 401
                               58.671 46.290 26.095 1.00 32.26
15
     ATOM 1869 CA PHE 401
                                59.899 45.682 26.596 1.00 35.15
     ATOM 1870 CB PHE 401
                                61.014 46.739 26.648 1.00 35.99
     ATOM 1871 CG PHE 401
                                62.346 46.213 27.117 1.00 39.41
                                62.845 45.003 26.639 1.00 35.94
     ATOM 1872 CD1 PHE 401
20
     ATOM 1873 CD2 PHE 401
                                63.119 46.944 28.019 1.00 40.55
                                64.088 44.531 27.055 1.00 30.16
     ATOM 1874 CE1 PHE 401
                                64.367 46.478 28.439 1.00 35.53
     ATOM 1875 CE2 PHE 401
                                64.849 45.271 27.952 1.00 36.39
     ATOM 1876 CZ PHE 401
                               59.607 45.129 27.996 1.00 36.42
     ATOM 1877 C PHE 401
                               59.957 43.995 28.317 1.00 36.71
25
    ATOM 1878 O PHE 401
                               58.920 45.925 28.805 1.00 36.59
     ATOM 1879 N LEU 402
     ATOM 1880 CA LEU 402
                                58.561 45.528 30.158 1.00 37.68
     ATOM 1881 CB LEU 402
                                57.986 46.720 30.917 1.00 40.71
     ATOM 1882 CG LEU 402
                                58.963 47.751 31.463 1.00 43.13
30
     ATOM 1883 CD1 LEU 402
                                58.180 48.926 32.031 1.00 39.88
     ATOM 1884 CD2 LEU 402
                                59.847 47.103 32.527 1.00 38.39
     ATOM 1885 C LEU 402
                               57.521 44.420 30.164 1.00 38.02
     ATOM 1886 O LEU 402
                               57.582 43.507 30.984 1.00 37.39
     ATOM 1887 N GLU 403
                               56.558 44.522 29.251 1.00 39.74
    ATOM 1888 CA GLU 403
                                55.469 43.559 29.166 1.00 42.79
35
     ATOM 1889 CB GLU 403
                                54.445 44.022 28.129 1.00 46.21
                                53.092 43.330 28.232 1.00 56.88
     ATOM 1890 CG GLU 403
                                52.090 43.833 27.202 1.00 65.21
     ATOM 1891 CD GLU 403
                                52.230 44.983 26.728 1.00 70.60
     ATOM 1892 OE1 GLU 403
                                51.154 43.073 26.870 1.00 70.53
    ATOM 1893 OE2 GLU 403
                               55,890 42.121 28.886 1.00 40.14
     ATOM 1894 C GLU 403
     ATOM 1895 O GLU 403
                               55.368 41.200 29.506 1.00 40.57
                               56.835 41.932 27.966 1.00 39.43
     ATOM 1896 N VAL 404
     ATOM 1897 CA VAL 404
                                57.292 40.586 27.610 1.00 40.96
     ATOM 1898 CB VAL 404
                                57.851 40.516 26.159 1.00 35.50
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     ATOM 1899 CG1 VAL 404
                                56.807 40.995 25.177 1.00 43.46
     ATOM 1900 CG2 VAL 404
                                59.132 41.321 26.030 1.00 25.74
     ATOM 1901 C VAL 404
                               58.317 39.946 28.536 1.00 41.94
     ATOM 1902 O VAL 404
                               58.468 38.722 28.533 1.00 43.82
                               59.026 40.759 29.310 1.00 39.84
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     ATOM 1903 N PHE 405
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60.051 40.223 30.189 1.00 42.73
    ATOM 1904 CA PHE 405
                                61.401 40.897 29.913 1.00 36.85
    ATOM 1905 CB PHE 405
                                61.963 40.596 28.551 1.00 33.23
    ATOM 1906 CG PHE 405
    ATOM 1907 CD1 PHE 405
                                 62.283 41.625 27.672 1.00 33.90
                                 62.157 39.281 28.138 1.00 31.62
    ATOM 1908 CD2 PHE 405
                                62.786 41.351 26.399 1.00 39.16
    ATOM 1909 CE1 PHE 405
                                62.657 38.997 26.872 1.00 33.33
    ATOM 1910 CE2 PHE 405
                                62.972 40.033 25.999 1.00 31.99
    ATOM 1911 CZ PHE 405
                               59,723 40,273 31,676 1,00 43,97
    ATOM 1912 C PHE 405
                               60.636 39.943 32.460 1.00 46.56
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    ATOM 1913 O PHE 405
                               67.928 36.755 11.188 1.00 33.04
    ATOM
              1 O1 HOH 501
                               69.618 40.719 13.009 1.00 23.00
    ATOM
             2 O1 HOH 502
    ATOM
             3 O1 HOH 503
                               64.885 40.168 12.340 1.00 23.00
                               63.079 40.108 15.841 1.00 23.00
             4 O1 HOH 504
    ATOM
                               63.404 46.536 15.354 1.00 36.41
    ATOM
             5 O1 HOH 505
15
    ATOM
             6 O1 HOH 506
                               61.299 15.617 -0.595 1.00 23.00
             7 O1 HOH 507
                               67.359 15.375 0.551 1.00 23.00
    ATOM
             8 O1 HOH 508
                               67,230 12.002 -0.634 1.00 23.00
    ATOM
             9 O1 HOH 509
                               66.906 12.467 3.855 1.00 23.00
     ATOM
                               61.785 9.946 3.983 1.00 23.00
             10 O1 HOH 510
20
     ATOM
             11 O1 HOH 511
                               57.670 11.385 9.909 1.00 23.00
     ATOM
             12 O1 HOH 512
                               55.791 11.570 10.291 1.00 23.00
    ATOM
             13 O1 HOH 513
                               54.637 14.058 9.201 1.00 23.00
    ATOM
                               55.882 16.054 12.204 1.00 26.53
             14 O1 HOH 514
     ATOM
                               53.685 15.842 18.209 1.00 23.00
             15 O1 HOH 515
25
     ATOM
             16 O1 HOH 516
                               49.559 24.773 19.020 1.00 23.00
     ATOM
                                51.258 25.512 13.384 1.00 37.74
             17 O1 HOH 517
     ATOM
             18 O1 HOH 518
                                53.551 25.749 10.593 1.00 42.31
     ATOM
                                50.338 23.299 7.662 1.00 41.19
             19 O1 HOH 519
     ATOM
             20 O1 HOH 520
                                50.830 20.272 8.323 1.00 28.46
     ATOM
30
                                48.630 20.291 6.429 1.00 23.00
             21 O1 HOH 521
     ATOM
             22 O1 HOH 522
                                49.233 17.389 2.867 1.00 23.00
     ATOM
             23 O1 HOH 523
                                52.076 22.770 1.260 1.00 23.00
     ATOM
             24 O1 HOH 524
                                51.671 23.621 -1.020 1.00 23.00
     ATOM
                                58.294 31.509 2.147 1.00 31.83
             25 O1 HOH 525
35
     ATOM
                                57.497 36.071 2.268 1.00 23.00
             26 O1 HOH 526
     ATOM
                                65.373 36.025 6.809 1.00 23.00
             27 O1 HOH 527
     ATOM
                                67.871 36.399 6.419 1.00 66.52
             28 O1 HOH 528
     ATOM
                                67.189 33.811 9.409 1.00 23.00
             29 O1 HOH 529
     ATOM
             30 O1 HOH 530
                                62.458 48.056 13.590 1.00 23.00
40
     ATOM
             31 O1 HOH 531
                                63.943 46.824 10.638 1.00 39.26
     ATOM
                                57.465 45.867 13.186 1.00 23.00
             32 O1 HOH 532
     ATOM
             33 O1 HOH 533
                                55.223 40.774 10.959 1.00 23.00
     ATOM
             34 O1 HOH 534
                                53.737 44.032 19.560 1.00 23.00
     ATOM
                                55.982 49.757 24.168 1.00 23.00
45
     ATOM
             35 O1 HOH 535
                                58.575 52.330 31.881 1.00 23.00
     ATOM
             36 O1 HOH 536
             37 O1 HOH 537
                                62.563 49.327 37.804 1.00 23.00
     ATOM
             38 O1 HOH 538
     ATOM
                                61.736 40.280 35.059 1.00 60.53
                                63.271 38.155 34.156 1.00 52.21
     ATOM
             39 O1 HOH 539
             40 O1 HOH 540
                                61.872 35.187 29.990 1.00 23.00
50
     ATOM
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	ATON 4	41 01 11011	E 4.1	63.701 36.808 28.720 1.00 23.00
	ATOM	41 O1 HOH	541	
	ATOM	42 O1 HOH	542	
	ATOM	43 O1 HOH	543	
_	ATOM	44 O1 HOH	544	-
5	ATOM	45 O1 HOH	545	
	ATOM	46 O1 HOH	546	<b>***</b>
	ATOM	47 O1 HOH	547	
	ATOM	48 O1 HOH	548	• • •
• •	ATOM	49 O1 HOH	549	
10	ATOM	50 O1 HOH	550	<b>32.</b> , , , , , , , , , , , , , , , , , , ,
	ATOM	51 O1 HOH	551	
	ATOM	52 O1 HOH	552	
	ATOM	53 O1 HOH	553	
1.5	ATOM	54 O1 HOH	554	
15	ATOM	55 O1 HOH	555	
	ATOM	56 O1 HOH	556	
	ATOM	57 O1 HOH	557	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -
	ATOM	58 O1 HOH	558	
20	ATOM	59 O1 HOH	559	
20	ATOM	60 O1 HOH	560	
	ATOM	61 O1 HOH	561	
	ATOM	62 O1 HOH	562	
	ATOM	63 O1 HOH	563 564	
25	ATOM	64 O1 HOH		
25	ATOM	65 O1 HOH 66 O1 HOH	565 566	76.525 41.395 10.460 1.00 23.00 79.358 49.535 15.048 1.00 53.78
	ATOM		567	78.046 53.530 9.188 1.00 23.00
	ATOM ATOM	67 O1 HOH 68 O1 HOH	568	68.058 52.158 15.548 1.00 23.00
	ATOM	69 O1 HOH	569	68.598 53.164 18.083 1.00 45.72
30	ATOM	70 O1 HOH	570	73.482 58.914 21.552 1.00 58.99
30	ATOM	71 O1 HOH	571	65.648 53.551 26.240 1.00 23.00
	ATOM	72 O1 HOH	572	75.776 46.207 30.367 1.00 33.32
	ATOM	72 O1 HOH	573	78.686 46.470 31.087 1.00 23.00
	ATOM	74 O1 HOH	574	77.580 41.209 31.884 1.00 23.00
35	ATOM	75 O1 HOH	575	76.879 31.531 24.067 1.00 23.00
55	ATOM	76 O1 HOH	576	77.927 29.163 20.647 1.00 23.00
	ATOM	77 O1 HOH	577	80.180 24.963 17.233 1.00 53.36
	ATOM	78 O1 HOH	578	80.631 25.802 15.508 1.00 23.00
	ATOM	79 O1 HOH	579	82.104 22.566 14.156 1.00 23.00
40	ATOM	80 O1 HOH	580	76.954 22.077 18.425 1.00 46.50
10	ATOM	81 O1 HOH	581	86.619 37.903 16.945 1.00 47.66
	ATOM	82 O1 HOH	582	83.586 42.305 18.576 1.00 23.00
	ATOM	83 O1 HOH	583	83.481 45.262 19.526 1.00 23.00
	ATOM	84 O1 HOH	584	66.787 32.864 33.796 1.00 23.00
45	ATOM	85 O1 HOH	585	59,447 33.572 30.734 1.00 23.00
	ATOM	86 O1 HOH	586	57.013 32.278 31.125 1.00 23.00
	ATOM	87 O1 HOH	587	58.084 29.428 24.648 1.00 24.06
	ATOM	88 O1 HOH	588	52.774 25.054 32.650 1.00 57.81
	ATOM	89 O1 HOH	589	53.800 24.465 34.834 1.00 23.00
50	ATOM	90 O1 HOH	590	47.195 30.205 30.414 1.00 23.00

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91 O1 HOH 591
                               48.978 35.051 30.228 1.00 23.00
    ATOM
             92 O1 HOH 592
                               49.280 39.962 31.041 1.00 23.00
    ATOM
             93 O1 HOH 593
                               42.329 32.230 20.993 1.00 23.00
    ATOM
             94 O1 HOH 594
                               44.199 32.910 19.088 1.00 23.00
    ATOM
             95 O1 HOH 595
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    ATOM
                               48.971 31.296 14.022 1.00 23.00
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            99 O1 HOH 599
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           100 O1 HOH 600
    ATOM
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            102 O1 HOH 602
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                                47.359 19.644 28.494 1.00 41.57
            103 O1 HOH 603
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    ATOM 2300 C ACY 701
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     ATOM 2301 O ACY 701
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     ATOM 2302 OXT ACY 701
    ATOM 2303 CH3 ACY 701
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                              67.309 42.207 18.510 1.00 32.20
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                         - 1
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     ATOM 2305 C2 IBR
                          1
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     ATOM 2306 C3 IBR
     ATOM 2307 C4 IBR
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                          1
                              67.884 43.772 20.218 1.00 35.08
     ATOM 2308 C5 IBR
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     ATOM 2309 C6 IBR
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                              68.673 42.828 20.790 1.00 30.76
     ATOM 2310 C7 IBR
                              67.681 43.327 25.704 1.00 29.18
     ATOM 2311 C8 IBR
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     ATOM 2312 C9 IBR
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     ATOM 2313 C10 IBR
     ATOM 2314 C11 IBR
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                           1
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     ATOM 2315 C12 IBR
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     ATOM 2316 C13 IBR
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     ATOM 2317 C14 IBR
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     ATOM 2318 C15 IBR
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     ATOM 2319 C16 IBR
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     ATOM 2320 C17 IBR
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     ATOM 2325 O1 IBR
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     ATOM 2326 O2 IBR
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     ATOM 2327 O3 IBR
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                           1
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     ATOM 2328 O4 IBR
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     END
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WO 99/26966

WO 99/26966

PCT/US98/25296

#### APPENDIX 6

### TR T3.PDB

REMARK rTR\_t3 full length numbering

**REMARK** 

5 REMARK Rfactor 0.221 Rfree 0.240

REMARK Resolution 5. 2.0 all reflections

REMARK conformation of MET 388 confirmed by SA\_omit map

REMARK

REMARK Three cacodylate-modified cysteines (CYA)

REMARK Cya334, Cya380, Cya392 10

REMARK cacodylate modeled as single arsenic atom

REMARK side chain of certain residues modeled as ALA due to poor density;

REMARK however, residue name reflects true residue for clarity

15 REMARK

REMARK clone obtained from Murray et. al.

REMARK deposited sequence confirmed,

REMARK differing from that reported by Thompson et. al.

REMARK in the following codons:

REMARK 281 Thr - Ala 20

REMARK 285 Lys - Glu

REMARK identical to that reported by Mitsuhashi et. al.

REMARK gb:RNTRAVI X07409

M.B. MURRAY, N.D.ZILZ, AUTH JRNL

25 N.L.MCCREARY, M.J.MACDONALD

> **AUTH 2 H.C.TOWLE** JRNL

TITL ISOLATION AND CHARACTERIZATION OF RAT CDNA **JRNL CLONES FOR TWO** 

TITL 2 DISTINCT THYROID HORMONE RECPTORS JRNL

V. 263 25 1988 30 JRNL REF JBC

AUTH C.C.THOMPSON, C.WEINBERGER, R.LEBO, R.M.EVANS JRNL IDENTIFICATION OF A NOVEL THYROID HORMONE JRNL TITL

RECEPTOR EXPRESSED

TITL 2 IN THE MAMMALIAN CENTRAL NERVOUS SYSTEM JRNL

35 **JRNL** REF SCIENCE

V. 237 1987 AUTH T.MITSUHASHI, G.TENNYSON, V.NIKODEM

**JRNL** JRNL TITL NUCLEOTIDE SEQUENCE OF NOVEL CDNAS GENERATED BY ALTERNATIVE

TITL 2 SPLICING OF A RAT THYROID HORMONE RECEPTOR **JRNL** 

GENE TRANSCRIPT 40

> JRNL REF NUC. ACIDS. RES. V. 16 12 1988

REMARK

45

68.406 10.620 7.027 1.00 41.66 ATOM 1 CB ARG 157

69.926 10.540 6.997 1.00 44.48 2 CG ARG 157 ATOM 70.552 11.261 8.173 1.00 47.02 3 CD ARG 157 ATOM

70.112 10.680 9.435 1.00 49.73 ATOM 4 NE ARG 157

ATOM 5 CZ ARG 157 70.917 10.392 10.450 1.00 51.21

6 NH1 ARG 157 72,223 10.629 10.361 1.00 51.79 ATOM

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ATOM
              7 NH2 ARG 157
                                70.405 9.871 11.556 1.00 51.92
     ATOM
              8 C ARG 157
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              9 O ARG 157
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     ATOM
             10 N ARG 157
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             11 CA ARG 157
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             13 CD PRO 158
             14 CA PRO 158
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     ATOM
             15 CB PRO 158
                               63.219 10.015 7.694 1.00 31.87
     ATOM
                               64.260 10.158 8.759 1.00 33.55
             16 CG PRO 158
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     ATOM
             17 C PRO 158
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             22 CG GLU 159
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     ATOM
             23 CD GLU 159
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             27 O GLU 159
             28 N PRO 160
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     ATOM
                               61.997 16.139 4.207 1.00 16.57
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             29 CD PRO 160
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             30 CA PRO 160
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             31 CB PRO 160
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             33 C PRO 160
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             38 OG1 THR 161
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PCT/US98/25296

## WO 99/26966

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_	ATOM	60 CB GLU 164	56.880 17.340 7.664 1.00 21.23
5	ATOM	61 CG GLU 164	56.509 16.508 8.886 1.00 20.30
	ATOM	62 CD GLU 164	57.557 15.483 9.243 1.00 20.07
	ATOM	63 OE1 GLU 164	58.409 15.186 8.385 1.00 19.80
	ATOM	64 OE2 GLU 164	57.532 14.977 10.385 1.00 21.00
	ATOM	65 C GLU 164	56.195 19.289 6.235 1.00 22.45
10	ATOM	66 O GLU 164	56.607 20.354 6.684 1.00 23.36
	ATOM	67 N TRP 165	56.140 19.024 4.928 1.00 21.06
	ATOM	68 CA TRP 165	56.518 20.031 3.936 1.00 19.57
	ATOM	69 CB TRP 165	56.486 19.466 2.518 1.00 16.06
	ATOM	70 CG TRP 165	57.775 18.839 2.120 1.00 14.01
15	ATOM	71 CD2 TRP 165	59.055 19.480 2.037 1.00 13.26
	ATOM	72 CE2 TRP 165	59.976 18.515 1.588 1.00 12.91
	ATOM	73 CE3 TRP 165	59.507 20.779 2.300 1.00 14.44
	ATOM	74 CD1 TRP 165	57.972 17.544 1.738 1.00 12.89
	ATOM	75 NEI TRP 165	59.290 17.343 1.413 1.00 12.80
20	ATOM	76 CZ2 TRP 165	61.328 18.805 1.388 1.00 15.06
	ATOM	77 CZ3 TRP 165	60.850 21.069 2.103 1.00 14.72
	ATOM	78 CH2 TRP 165	61.747 20.084 1.649 1.00 16.82
	ATOM	79 C TRP 165	55.553 21.210 4.056 1.00 18.93
	ATOM	80 O TRP 165	55.960 22.359 3.926 1.00 21.12
25	ATOM	81 N ASP 166	54.279 20.922 4.307 1.00 19.33
	ATOM	82 CA ASP 166	53.262 21.963 4.483 1.00 20.35
	ATOM	83 CB ASP 166	51.864 21.353 4.672 1.00 20.22
	ATOM	84 CG ASP 166	51.302 20.748 3.386 1.00 23.36
	ATOM	85 OD1 ASP 166	51.746 21.153 2.296 1.00 23.42
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	ATOM	87 C ASP 166	53.623 22.785 5.712 1.00 21.02
	ATOM	88 O ASP 166	53.627 24.013 5.654 1.00 22.56
	ATOM	89 N LEU 167	53.926 22.096 6.813 1.00 20.50
	ATOM	90 CA LEU 167	54.312 22.726 8.071 1.00 21.37
35	ATOM	91 CB LEU 167	54.661 21.657 9.109 1.00 23.49
	ATOM	92 CG LEU 167	54.223 21.846 10.565 1.00 27.19
	ATOM	93 CD1 LEU 167	55.312 21.291 11.453 1.00 27.70
	ATOM	94 CD2 LEU 167	53.940 23.314 10.906 1.00 27.71
	ATOM	95 C LEU 167	55.541 23.602 7.839 1.00 20.72
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	ATOM		56.505 23.051 7.114 1.00 18.54
	ATOM	98 CA ILE 168	57.747 23.725 6.778 1.00 18.60
	ATOM	99 CB ILE 168	58.671 22.771 5.995 1.00 17.54
	ATOM	100 CG2 ILE 168	59.695 23.533 5.163 1.00 17.65
45	ATOM	101 CG1 ILE 168	59.330 21.794 6.972 1.00 20.27
	ATOM	102 CD1 ILE 168	60.048 20.631 6.322 1.00 17.96
	ATOM	103 C ILE 168	57.486 25.002 5.979 1.00 21.96
	ATOM	104 O ILE 168	58.045 26.064 6.291 1.00 23.06
	ATOM	105 N HIS 169	56.591 24.925 4.996 1.00 22.04
50	ATOM	106 CA HIS 169	56.285 26.092 4.164 1.00 21.21

WO 99/26966

PCT/US98/25296

	ATOM	107 CB HIS 169	55.413 25.702 2.969 1.00 20.12
	ATOM	108 CG HIS 169	56.101 24.799 2.001 1.00 19.18
	ATOM	109 CD2 HIS 169	57.398 24.733 1.619 1.00 18.62
	ATOM	110 ND1 HIS 169	55.457 23.764 1.357 1.00 17.90
5	ATOM	111 CE1 HIS 169	56.327 23.096 0.625 1.00 18.43
	ATOM	112 NE2 HIS 169	57.513 23.660 0.772 1.00 20.10
	ATOM	113 C HIS 169	55.615 27.198 4.959 1.00 20.61
	ATOM	114 O HIS 169	55.979 28.370 4.836 1.00 20.08
	ATOM	115 N VAL 170	54.632 26.821 5.769 1.00 20.01
10	ATOM	116 CA VAL 170	53.922 27.785 6.580 1.00 20.52
	ATOM	117 CB VAL 170	52.816 27.120 7.384 1.00 21.33
	ATOM	118 CG1 VAL 170	
	ATOM	119 CG2 VAL 170	
	ATOM	120 C VAL 170	54.891 28.477 7.521 1.00 20.58
15	ATOM	121 O VAL 170	54.926 29.704 7.554 1.00 22.32
	ATOM	122 N ALA 171	55.712 27.696 8.230 1.00 18.83
	ATOM	123 CA ALA 171	56.692 28.234 9.182 1.00 18.34
	ATOM	124 CB ALA 171	57.375 27.102 9.946 1.00 17.05
	ATOM	125 C ALA 171	57.733 29.151 8.533 1.00 17.84
20	ATOM	126 O ALA 171	58.084 30.200 9.091 1.00 18.67
20	ATOM	127 N THR 172	58.231 28.756 7.367 1.00 17.81
	ATOM	128 CA THR 172	59.215 29.551 6.639 1.00 18.88
	ATOM	129 CB THR 172	59.726 28.794 5.380 1.00 20.47
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25	ATOM	131 CG2 THR 172	
	ATOM	132 C THR 172	58.655 30.932 6.251 1.00 19.42
	ATOM	133 O THR 172	59.320 31.957 6.435 1.00 17.98
	ATOM	134 N GLU 173	57.425 30.970 5.756 1.00 19.97
	ATOM	135 CA GLU 173	56.811 32.236 5.374 1.00 22.51
30	ATOM	136 CB GLU 173	55.520 31.981 4.577 1.00 27.26
20	ATOM	137 CG GLU 173	54.823 33.244 4.005 1.00 34.96
	ATOM	138 CD GLU 173	55.690 34.040 3.020 1.00 39.54
	ATOM	139 OE1 GLU 173	56.610 33.454 2.395 1.00 41.82
	ATOM	140 OE2 GLU 173	55.443 35.259 2.872 1.00 41.06
35	ATOM	141 C GLU 173	56.538 33.099 6.622 1.00 21.60
33	ATOM	142 O GLU 173	56.726 34.313 6.595 1.00 21.73
	ATOM	143 N ALA 174	56.123 32.461 7.716 1.00 19.69
	ATOM		55.844 33.155 8.968 1.00 18.07
	ATOM		55.423 32.169 10.037 1.00 16.90
40	ATOM	146 C ALA 174	
	ATOM	147 O ALA 174	
	ATOM	148 N HIS 175	58.240 33.222 9.259 1.00 16.39
	ATOM	149 CA HIS 175	59.498 33.831 9.629 1.00 16.41
	ATOM	150 CB HIS 175	60.574 32.758 9.804 1.00 12.71
45	ATOM	150 CB HIS 175	61.938 33.318 10.043 1.00 11.09
73	ATOM	151 CO 1113 175	62.373 34.252 10.920 1.00 8.26
	ATOM	152 CD2 HIS 175 153 ND1 HIS 175	63.030 32.977 9.273 1.00 13.39
	ATOM	154 CE1 HIS 175	63.030 32.977 9.273 1.00 13.39 64.076 33.683 9.658 1.00 13.77
	ATOM		63.702 34.464 10.658 1.00 12.70
50	ATOM		59.959 34.903 8.624 1.00 19.55
20	ATOM	156 C HIS 175	0.024 1.00 17.33

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            158 N ARG 176
    ATOM
                               60.424 35.494 6.307 1.00 21.30
    ATOM
            159 CA ARG 176
                               60.315 34.876 4.917 1.00 24.87
    ATOM
            160 CB ARG 176
                               61.361 33.827 4.609 1.00 30.22
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            161 CG ARG 176
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            162 CD ARG 176
    ATOM
            163 NE ARG 176
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            164 CZ ARG 176
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                                61.000 31.935 0.894 1.00 50.83
            165 NH1 ARG 176
    ATOM
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            166 NH2 ARG 176
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    ATOM
                              59.658 36.807 6.337 1.00 20.67
    ATOM
            167 C ARG 176
                              60.256 37.877 6.238 1.00 20.53
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            168 O ARG 176
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            169 N SER 177
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            171 CB SER 177
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            172 OG SER 177
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            174 O SER 177
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            175 N THR 178
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            177 CB THR 178
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            181 O THR 178
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            182 N ASN 179
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            186 OD1 ASN 179
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            189 O ASN 179
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            191 CA ALA 180
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    ATOM
            193 C ALA 180
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            194 O ALA 180
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            201 NE2 GLN 181
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PCT/US98/25296

WO 99/26966

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            208 N SER 183
                              66.926 39.272 3.224 1.00 43.72
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                               68.299 39.001 2.812 1.00 45.88
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            209 CA SER 183
                               68.304 38.069 1.593 1.00 47.26
     ATOM
            210 CB SER 183
            211 OG SER 183
                               67.519 38.605 0.531 1.00 47.23
     ATOM
            212 C SER 183
                              69.095 40.268 2.497 1.00 46.24
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                              70.290 40.194 2.185 1.00 48.13
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            213 O SER 183
                             68.445 41.426 2.579 1.00 45.79
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                             69.111 42.690 2.276 1.00 45.00
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                             68.127 43.636 1.594 1.00 43.54
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                             69.732 43.351 3.516 1.00 44.67
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            217 C HIS 184
                              70.316 44.440 3.428 1.00 45.02
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            218 O HIS 184
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            219 N TRP 185
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            220 CA TRP 185
     ATOM
            221 CB TRP 185
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            222 CG TRP 185
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     ATOM
                               72.197 40.593 7.291 1.00 33.38
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            223 CD2 TRP 185
                               72.572 39.321 6.807 1.00 31.68
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            224 CE2 TRP 185
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            225 CE3 TRP 185
                               73.092 41.296 8.107 1.00 31.65
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            226 CD1 TRP 185
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            227 NE1 TRP 185
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                               73.795 38.733 7.121 1.00 31.67
            228 CZ2 TRP 185
    ATOM
     ATOM
            229 CZ3 TRP 185
                               74.308 40.713 8.419 1.00 31.29
     ATOM
            230 CH2 TRP 185
                               74.651 39.444 7.923 1.00 31.06
            231 C TRP 185
                              71.618 43.720 5.856 1.00 41.52
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     ATOM
                              71.893 44.817 6.335 1.00 40.52
     ATOM
            232 O TRP 185
            233 N LYS 186
                              72.520 42.976 5.234 1.00 42.94
     ATOM
            234 CA LYS 186
                               73.896 43.417 5.143 1.00 45.25
    ATOM
            235 CB LYS 186
                               74.764 42.328 4.508 1.00 45.96
     ATOM
     ATOM
            236 CG LYS 186
                               76.255 42.600 4.590 1.00 48.07
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    ATOM
            237 CD LYS 186
                               77.053 41.307 4.504 1.00 51.20
                               78.554 41.574 4.457 1.00 52.69
    ATOM
            238 CE LYS 186
                               78.975 42.277 3.201 1.00 55.56
    ATOM
            239 NZ LYS 186
                              74.025 44.730 4.377 1.00 47.38
     ATOM
            240 C LYS 186
            241 O LYS 186
                              74.914 45.535 4.663 1.00 47.65
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     ATOM
                              73.134 44.959 3.418 0.50 48.02
            242 N GLN 187
    ATOM
                                                             ALTA
            243 CA GLN 187
                               73.193 46.183 2.623 0.50 48.69
                                                             ALTA
    ATOM
    ATOM
            244 CB GLN 187
                               72.547 45.973 1.246 0.50 48.66
                                                             ALTA
                               73.104 44.771 0.453 0.50 49.05
                                                             ALTA
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            245 CG GLN 187
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            246 CD GLN 187
                               74.624 44.766 0.339 0.50 49.17
                                                             ALTA
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            247 OE1 GLN 187
                               75,225 45.691 -0.209 0.50 49.71
                                                             ALTA
                               75.250 43.710 0.847 0.50 48.57
            248 NE2 GLN 187
                                                            ALTA
     ATOM
                              72.551 47.373 3.343 0.50 49.06
    ATOM
            249 C GLN 187
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                               73.094 48.475 3.329 0.50 49.53
     ATOM
            250 O GLN 187
                                                             ALTA
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            251 N ARG 188
                               71.405 47.152 3.980 1.00 49.18
    ATOM
                               70.723 48.221 4.695 1.00 49.90
    ATOM
            252 CA ARG 188
                               69.209 47.988 4.653 1.00 53.68
    ATOM
            253 CB ARG 188
                               68.617 47.798 3.251 1.00 57.22
    ATOM 254 CG ARG 188
                               67.099 47.962 3.302 1.00 60.67
            255 CD ARG 188
    ATOM
                               66.430 47.441 2.110 1.00 64.43
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    ATOM 256 NE ARG 188
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65.931 46.208 2.009 1.00 66.13
            257 CZ ARG 188
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                                66.027 45.362 3.031 1.00 66.69
            258 NH1 ARG 188
    ATOM
                                65.318 45.823 0.893 1.00 66.10
            259 NH2 ARG 188
    ATOM
                               71.150 48.510 6.133 1.00 48.42
            260 C ARG 188
    ATOM
                               70.544 49.368 6.784 1.00 48.86
    ATOM
            261 O ARG 188
                               72.153 47.804 6.647 1.00 46.00
    ATOM
            262 N ARG 189
                                72.581 48.030 8.028 1.00 44.24
    ATOM
            263 CA ARG 189
                                73.039 46.726 8.690 1.00 43.40
    ATOM
            264 CB ARG 189
                                74.367 46.204 8.203 1.00 43.05
            265 CG ARG 189
    ATOM
                                74.808 45.021 9.019 1.00 43.62
    ATOM
            266 CD ARG 189
10
                                76.185 44.660 8.717 1.00 45.95
    ATOM
            267 NE ARG 189
                                76.981 43.976 9.536 1.00 48.56
    ATOM
            268 CZ ARG 189
            269 NH1 ARG 189
                                76.548 43.560 10.724 1.00 46.34
    ATOM
                                78.233 43.735 9.174 1.00 50.12
            270 NH2 ARG 189
    ATOM
                               73.642 49.116 8.238 1.00 43.20
            271 C ARG 189
    ATOM
15
            272 O ARG 189
                               74.629 49.210 7.500 1.00 43.07
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            273 N LYS 190
                               73.427 49.925 9.268 1.00 41.56
    ATOM
                               74.335 51.003 9.628 1.00 39.96
            274 CA LYS 190
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                               73.563 52.323 9.757 1.00 38.85
    ATOM
           275 CB LYS 190
                               74.983 50.631 10.956 1.00 38.91
           276 C LYS 190
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    ATOM
            277 O LYS 190
                               74.345 50.015 11.806 1.00 38.17
    ATOM
                               76.261 50.959 11.104 1.00 38.49
            278 N PHE 191
    ATOM
                               76.998 50.673 12.326 1.00 38.42
    ATOM
            279 CA PHE 191
                               78.500 50.762 12.073 1.00 38.37
            280 CB PHE 191
     ATOM
    ATOM
                                79.056 49.608 11.308 1.00 39.05
25
            281 CG PHE 191
                                78.712 49.408 9.976 1.00 40.02
            282 CD1 PHE 191
    ATOM
                                79.942 48.727 11.917 1.00 39.19
            283 CD2 PHE 191
    ATOM
                                79.245 48.344 9.256 1.00 40.57
            284 CE1 PHE 191
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                                80.482 47.661 11.213 1.00 40.32
            285 CE2 PHE 191
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                               80.133 47.466 9.875 1.00 41.84
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                               76.433 51.184 14.634 1.00 37.05
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            290 CA LEU 192
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                                75.833 51.247 17.014 1.00 33.04
            291 CB LEU 192
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                                75.503 52.074 18.260 1.00 31.38
            292 CG LEU 192
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                                74.116 52.651 18.102 1.00 29.02
            293 CD1 LEU 192
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                                75.592 51.229 19.536 1.00 30.32
            294 CD2 LEU 192
     ATOM
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                               78,500 52.218 16.112 1.00 37.66
            296 O LEU 192
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     ATOM
                               77.377 54.177 15.988 1.00 38.15
             297 N PRO 193
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                                76.156 54.996 15.902 1.00 37.90
     ATOM
            298 CD PRO 193
                                78.561 55.025 16.187 1.00 38.68
            299 CA PRO 193
     ATOM
            300 CB PRO 193
                                77.950 56.365 16.568 1.00 37.20
     ATOM
                                76.711 56.397 15.758 1.00 37.08
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     ATOM
            301 CG PRO 193
                               79.475 54.503 17.294 1.00 41.12
     ATOM
            302 C PRO 193
                               79.005 54.129 18.367 1.00 42.26
             303 O PRO 193
     ATOM
                               80.782 54.509 17.052 1.00 43.62
     ATOM
             304 N ASP 194
                               81.731 54.012 18.050 1.00 46.71
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             305 CA ASP 194
            306 CB ASP 194
                                83.131 53.938 17.470 1.00 49.32
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307 CG ASP 194
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                                83.539 51.726 16.719 1.00 53.18
            308 OD1 ASP 194
    ATOM
                                82.981 53.268 15.227 1.00 55.10
            309 OD2 ASP 194
    ATOM
                              81.769 54.743 19.386 1.00 47.12
            310 C ASP 194
    ATOM
                              82.158 54.163 20.403 1.00 48.16
            311 O ASP 194
    ATOM
                              81.389 56.015 19.386 1.00 47.54
            312 N ASP 195
    ATOM
                               81.382 56.791 20.620 1.00 48.68
    ATOM 313 CA ASP 195
                               81.180 58.285 20.322 1.00 50.76
            314 CB ASP 195
    ATOM
                               79.871 58.572 19.602 1.00 54.24
            315 CG ASP 195
    ATOM
                                78.929 59.082 20.253 1.00 56.17
            316 OD1 ASP 195
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    ATOM
                                79.786 58.292 18.385 1.00 56.08
            317 OD2 ASP 195
    ATOM
                              80,304 56,274 21.580 1.00 47.63
            318 C ASP 195
    ATOM
                               80.294 56.621 22.772 1.00 49.07
    ATOM
            319 O ASP 195
            320 N ILE 196
                              79.400 55.444 21.065 1.00 44.87
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            321 CA ILE 196
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                               76.983 54.813 21.121 1.00 42.19
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            322 CB ILE 196
                               75.870 54.357 22.060 1.00 40.29
    ATOM
            323 CG2 ILE 196
                               76.635 56.191 20.535 1.00 41.32
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     ATOM
                               75.344 56.219 19.732 1.00 41.32
            325 CD1 ILE 196
    ATOM
    ATOM 326 C ILE 196
                              78.725 53.509 22.391 1.00 40.89
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     ATOM 327 O ILE 196
                              79.358 52.722 21.679 1.00 40.08
            328 N GLY 197
                               78.384 53.240 23.642 1.00 40.16
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                                78.705 51.957 24.228 1.00 40.21
            329 CA GLY 197
     ATOM
            330 C GLY 197
                               80.066 51.907 24.879 1.00 40.18
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            331 O GLY 197
                               80.512 50.839 25.267 1.00 40.55
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     ATOM
            332 N GLN 198
                               80,718 53.057 25.029 1.00 41.25
     ATOM
            333 CA GLN 198
                                82.038 53.111 25.664 1.00 40.94
     ATOM
            334 CB GLN 198
                                83.041 53.823 24.738 1.00 39.51
     ATOM
            335 C GLN 198
                               81.995 53.796 27.046 1.00 40.93
     ATOM
                               83.036 54.197 27.571 1.00 41.83
            336 O GLN 198
     ATOM
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            337 N SER 199
                               80.806 53.859 27.654 1.00 39.68
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                               80.615 54.510 28.961 1.00 37.74
            338 CA SER 199
     ATOM
                                79.995 55.905 28.768 1.00 38.50
            339 CB SER 199
     ATOM
                                80.687 56.672 27.792 1.00 40.71
            340 OG SER 199
     ATOM
                               79.743 53.726 29.958 1.00 36.31
            341 C SER 199
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     ATOM
                               78.719 54.228 30.436 1.00 35.69
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     ATOM
                               80.123. 52.484 30.280 1.00 35.05
            343 N PRO 200
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                                81.246 51.684 29.760 1.00 33.97
            344 CD PRO 200
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                                79.313 51.715 31.228 1.00 35.89
            345 CA PRO 200
     ATOM
                                79.872 50.304 31.075 1.00 33.94
            346 CB PRO 200
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                                81,297 50,532 30,708 1,00 33,31
            347 CG PRO 200
     ATOM
            348 C PRO 200
                               79.477 52.241 32.656 1.00 37.75
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                               80.484 51.959 33.299 1.00 38.78
     ATOM
            349 O PRO 200
                              78.493 52.988 33.158 1.00 39.61
            350 N ILE 201
     ATOM
                               78.590 53.551 34.511 1.00 40.56
            351 CA ILE 201
45
     ATOM
                               78.715 55.093 34.484 1.00 40.20
     ATOM 352 CB ILE 201
                                80.125 55.501 34.082 1.00 41.06
     ATOM 353 CG2 ILE 201
                                77.690 55.694 33.532 1.00 40.98
     ATOM 354 CG1 ILE 201
                                77.969 57.147 33.205 1.00 44.31
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     ATOM
                              77.535 53.160 35.546 1.00 41.40
            356 C ILE 201
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	ATOM	359 CA VAL 202	75.325 52.293 36.053 1.00 40.70
	ATOM	360 CB VAL 202	73.913 52.292 35.422 1.00 38.44
5	ATOM	361 CG1 VAL 202	72.881 51.826 36.435 1.00 35.91
	ATOM	362 CG2 VAL 202	73.560 53.692 34.934 1.00 36.42
	ATOM	363 C VAL 202	75.687 50.917 36.622 1.00 41.64
	ATOM	364 O VAL 202	76.094 50.008 35.894 1.00 42.05
	ATOM	365 N SER 203	75.596 50.800 37.938 1.00 43.06
10	ATOM	366 CA SER 203	75.947 49.576 38.639 1.00 44.57
	ATOM	367 CB SER 203	75.916 49.842 40.154 1.00 46.82
	ATOM	368 OG SER 203	76,457 48.772 40.916 1.00 50.18
	ATOM	369 C SER 203	75.052 48.388 38.294 1.00 44.08
	ATOM	370 O SER 203	73.849 48.534 38.093 1.00 44.28
15	ATOM	371 N MET 204	75.656 47.210 38.231 1.00 43.11
15	ATOM	372 CA MET 204	74.930 45.980 37.963 1.00 43.12
	ATOM	373 CB MET 204	75.048 45.557 36.494 1.00 41.07
		374 CG MET 204	74.126 46.320 35.554 1.00 36.96
	ATOM ATOM	374 CG MET 204 375 SD MET 204	72.375 46.134 35.990 1.00 38.66
20		376 CE MET 204	71.970 44.592 35.098 1.00 37.26
20	ATOM		75.561 44.943 38.866 1.00 43.68
	ATOM	377 C MET 204	
	ATOM	378 O MET 204	76.784 44.817 38.912 1.00 44.32
	ATOM	379 N PRO 205	74.735 44.204 39.619 1.00 44.22
	ATOM	380 CD PRO 205	73.261 44.310 39.610 1.00 44.44
25	ATOM	381 CA PRO 205	75.187 43.164 40.546 1.00 44.32
	ATOM	382 CB PRO 205	73.944 42.299 40.701 1.00 45.18
	ATOM	383 CG PRO 205	72.832 43.335 40.691 1.00 44.29
	ATOM	384 C PRO 205	76.417 42.354 40.122 1.00 44.31
	ATOM	385 O PRO 205	77.393 42.293 40.864 1.00 43.97
30	ATOM	386 N ASP 206	76.404 41.802 38.912 1.00 44.30
	ATOM	387 CA ASP 206	77.524 40.984 38.433 1.00 44.77
	ATOM	388 CB ASP 206	77.073 40.106 37.270 1.00 47.12
	ATOM	389 CG ASP 206	76.503 40.912 36.120 1.00 49.73
	ATOM	390 OD1 ASP 206	76.992 42.039 35.863 1.00 49.65
35	ATOM	391 OD2 ASP 206	75.553 40.416 35.478 1.00 51.96
	ATOM	392 C ASP 206	78.805 41.718 38.037 1.00 44.10
	ATOM	393 O ASP 206	79.754 41.099 37.549 1.00 43.60
	ATOM	394 N GLY 207	78.804 43.039 38.145 1.00 44.19
	ATOM	395 CA GLY 207	80.001 43.785 37.803 1.00 43.51
40	ATOM	396 C GLY 207	80.041 44.425 36.433 1.00 43.29
	ATOM	397 O GLY 207	80.745 45.421 36.257 1.00 44.47
	ATOM	398 N ASP 208	79.363 43.845 35.446 1.00 42.45
	ATOM	399 CA ASP 208	79.347 44.436 34.106 1.00 41.51
	ATOM	400 CB ASP 208	78.915 43.402 33.070 1.00 42.91
45	ATOM	401 CG ASP 208	80.001 42.379 32.785 1.00 43.57
43	ATOM	401 CG ASP 208 402 OD1 ASP 208	79.675 41.218 32.468 1.00 44.55
		402 OD1 ASP 208 403 OD2 ASP 208	81.191 42.742 32.868 1.00 47.14
	ATOM		78,378 45.606 34.143 1.00 40.78
	ATOM	404 C ASP 208	77.176 45.403 34.277 1.00 42.50
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50	ATOM	406 N LYS 209	78.902 46.827 34.058 1.00 39.10

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407 CA LYS 209
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                               78.910 49.211 34.681 1.00 37.29
            408 CB LYS 209
     ATOM
            409 C LYS 209
                               77.326 48.423 32.871 1.00 34.47
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                               77.707 48.013 31.776 1.00 33.85
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                               76.275 49.228 33.028 1.00 33.30
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                                75.448 49.684 31.907 1.00 31.78
            412 CA VAL 210
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            413 CB VAL 210
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                                73.102 50.012 31.010 1.00 29.24
            414 CG1 VAL 210
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                                73.541 48.237 32.698 1.00 29.84
            415 CG2 VAL 210
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            416 C VAL 210
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            417 O VAL 210
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                               75.978 52.574 29.476 1.00 31.85
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            419 CA ASP 211
            420 CB ASP 211
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            421 CG ASP 211
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            422 OD1 ASP 211
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            425 O ASP 211
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            427 CA LEU 212
            428 CB LEU 212
                                72.440 55.736 30.470 1.00 32.41
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                                72.311 55.336 31.936 1.00 32.11
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            429 CG LEU 212
            430 CD1 LEU 212
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            431 CD2 LEU 212
            432 C LEU 212
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            433 O LEU 212
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                               73.370 55.589 27.407 1.00 32.21
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            435 CA GLU 213
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            436 CB GLU 213
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            437 CG GLU 213
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            438 CD GLU 213
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            439 OE1 GLU 213
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            440 OE2 GLU 213
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            441 C GLU 213
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            442 O GLU 213
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            443 N ALA 214
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                                73,769 52,585 24,482 1.00 30,43
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            446 C ALA 214
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            447 O ALA 214
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                                70.882 51.116 26.492 1.00 29.05
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     ATOM
            450 CB PHE 215
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                                69.443 50.689 28.535 1.00 25.53
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     ATOM
            451 CG PHE 215
                                69.330 49.344 28.854 1.00 26.16
     ATOM
            452 CD1 PHE 215
            453 CD2 PHE 215
                                68.349 51.519 28.737 1.00 25.04
     ATOM
                                68.144 48.831 29.370 1.00 25.73
     ATOM 454 CE1 PHE 215
                                67.160 51.018 29.252 1.00 25.84
            455 CE2 PHE 215
     ATOM
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     ATOM 456 CZ PHE 215
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             458 O PHE 215
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             459 N SER 216
                               69.714 53.108 25.776 1.00 31.41
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             460 CA SER 216
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            461 CB SER 216
                                68.976 55.375 25.256 1.00 32.50
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            462 OG SER 216
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PCT/US98/25296

## WO 99/26966

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	ATOM	577 OD2 ASP 231	54.443 41.784 13.522 1.00 25.90
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5	ATOM	711 CB ILE 248	50.473 35.824 27.273 1.00 24.59
	ATOM	712 CG2 ILE 248	51.304 36.682 28.242 1.00 24.09
	ATOM	713 CG1 ILE 248	49,499 36.707 26.487 1.00 23.47
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	ATOM	722 CD1 ILE 249	51.252 30.911 31.175 1.00 25.03
	ATOM	723 C ILE 249	54.481 32.148 27.470 1.00 22.24
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20	ATOM	726 CA LEU 250	55.028 31.079 25.345 1.00 21.40
	ATOM	727 CB LEU 250	54.319 30.290 24.239 1.00 20.06
	ATOM	728 CG LEU 250	53.566 29.038 24.677 1.00 20.22
	ATOM	729 CD1 LEU 250	52.952 28.406 23.453 1.00 19.19
	ATOM	730 CD2 LEU 250	54.494 28.050 25.386 1.00 18.52
25	ATOM	731 C LEU 250	55.850 32.209 24.736 1.00 20.82
	ATOM	732 O LEU 250	57.069 32.094 24.603 1.00 20.27
	ATOM	733 N LEU 251	55.179 33.302 24.384 1.00 22.14
	ATOM	734 CA LEU 251	55.842 34.467 23.805 1.00 22.90
	ATOM	735 CB LEU 251	54.806 35.543 23.471 1.00 22.76
30	ATOM	736 CG LEU 251	54.513 35.899 22.012 1.00 23.35
	ATOM	737 CD1 LEU 251	55.347 35.103 21.047 1.00 22.38
	ATOM	738 CD2 LEU 251	53.040 35.708 21.747 1.00 22.86
	ATOM	739 C LEU 251	56.891 35.030 24.776 1.00 23.67
	ATOM	740 O LEU 251	58.051 35.234 24.402 1.00 22.58
35	ATOM	741 N LYS 252	56.491 35.236 26.029 1.00 24.64
	ATOM	742 CA LYS 252	57.395 35.754 27.057 1.00 26.22
	ATOM	743 CB LYS 252	56.617 36.037 28.350 1.00 27.79
	ATOM	744 CG LYS 252	55.351 36.838 28.093 1.00 32.69
	ATOM	745 CD LYS 252	55.185 38.023 29.003 1.00 35.85
40	ATOM	746 CE LYS 252	54.773 37.626 30.397 1.00 39.34
	ATOM	747 NZ LYS 252	54.477 38.870 31.168 1.00 44.60
	ATOM	748 C LYS 252	58.566 34.793 27.312 1.00 25.26
	ATOM	749 O LYS 252	59.701 35.222 27.555 1.00 26.67
	ATOM	750 N GLY 253	58.306 33.497 27.195 1.00 23.97
45	ATOM	751 CA GLY 253	59.356 32.521 27.404 1.00 22.00
	ATOM	752 C GLY 253	60.397 32.429 26.292 1.00 23.10
	ATOM	753 O GLY 253	61.568 32.165 26.585 1.00 25.12
	ATOM	754 N CYS 254	60.014 32.702 25.041 1.00 22.27
	ATOM	755 CA CYS 254	60.944 32.584 23.908 1.00 20.91
50	ATOM	756 CB CYS 254	60.353 31.648 22.845 1.00 21.46

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58,992 32,385 21,893 1,00 22,92
             757 SG CYS 254
     ATOM
             758 C CYS 254
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     ATOM
                               62.215 33.834 22.316 1.00 19.88
             759 O CYS 254
     ATOM
                               60.731 34.984 23.561 1.00 19.56
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                                61.018 36.264 22.917 1.00 21.16
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    ATOM
                                60.292 37.407 23.634 1.00 21.21
            762 CB CYS 255
     ATOM
                                60.404 38.957 22.735 1.00 22.22
            763 SG CYS 255
     ATOM
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                                65.255 36.967 25.253 1.00 20.39
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             769 CG MET 256
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             770 SD MET 256
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     ATOM
             771 CE MET 256
                                68.856 38.971 24.375 1.00 18.47
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             772 C MET 256
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     ATOM
             773 O MET 256
                                66.305 36.164 22.225 1.00 18.15
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                                65.035 34.568 23.170 1.00 19.00
     ATOM
             774 N GLU 257
                                65.685 33.480 22.443 1.00 19.71
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             775 CA GLU 257
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     ATOM
            776 CB GLU 257
                                65.104 32.145 22.882 1.00 21.15
                                65.451 31.821 24.319 1.00 26.39
             777 CG GLU 257
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             778 CD GLU 257
                                64.513 30.820 24.929 1.00 30.75
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            779 OEI GLU 257
                                63.875 30.069 24.162 1.00 32.36
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            780 OE2 GLU 257
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     ATOM
                                66,521 33.506 20.197 1.00 17.58
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     ATOM
             783 N ILE 258
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             785 CB ILE 258
                               62.590 34.267 18.765 1.00 16.35
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     ATOM
             786 CG2 ILE 258
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30
                                61.935 32.884 18.980 1.00 17.24
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             788 CD1 ILE 258
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     ATOM
                              64.872 35.408 18.595 1.00 19.11
     ATOM
             789 C ILE 258
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             790 O ILE 258
                                64.785 36.517 19.341 1.00 19.71
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            792 CA MET 259
793 CB MET 259
                                65.486 37.744 18.956 1.00 18.43
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                                65.162 38.890 19.910 1.00 19.99
     ATOM
                                63,700 39,278 19,962 1,00 21,15
            794 CG MET 259
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             795 SD MET 259
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                                63.769 40.595 22.415 1.00 22.50
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             796 CE MET 259
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                                66.993 37.540 18.888 1.00 18.64
             797 C MET 259
     ATOM
             798 O MET 259
                                67.638 37.993 17.941 1.00 19.96
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             799 N SER 260
                               67.556 36.858 19.884 1.00 17.37
     ATOM
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             800 CA SER 260
                                69.387 35.840 21.195 1.00 17.25
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             801 CB SER 260
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             802 OG SER 260
             803 C SER 260
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             804 O SER 260
                               70.460 35.941 18.137 1.00 16.62
     ATOM
                               68.539 34.781 18.385 1.00 15.15
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     ATOM
             806 CA LEU 261
                                68.802 33.900 17.262 1.00 15.31
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             808 CG LEU 261
     ATOM
             809 CD1 LEU 261
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     ATOM
             810 CD2 LEU 261
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                               68.839 34.741 16.001 1.00 16.31
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             812 O LEU 261
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            817 CD ARG 262
     ATOM
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            818 NE ARG 262
     ATOM
                                61.755 36.932 13.361 1.00 21.06
            819 CZ ARG 262
     ATOM<sup>*</sup>
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             820 NH1 ARG 262
     ATOM
     ATOM
             821 NH2 ARG 262
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             822 C ARG 262
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    ATOM
             823 O ARG 262
             824 N ALA 263
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                               69.578 37.832 15.650 1.00 17.77
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             825 CA ALA 263
                                70.795 38.647 15.637 1.00 18.41
                                70.996 39.337 17.004 1.00 18.26
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    ATOM
                               71.998 37.740 15.327 1.00 19.15
             827 C ALA 263
     ATOM
             828 O ALA 263
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                               72.056 36.587 15.996 1.00 19.84
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                                73.155 35.633 15.818 1.00 20.35
             830 CA ALA 264
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            832 C ALA 264
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            833 O ALA 264
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             836 CB VAL 265
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                                 69.838 34.708 11.698 1.00 18.96
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            843 CB ARG 266
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            845 CD ARG 266
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                                67.833 39.940 9.069 1.00 23.26
     ATOM
            847 CZ ARG 266
     ATOM
             848 NH1 ARG 266
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     ATOM
             849 NH2 ARG 266
                                 66.960 39.733 8.099 1.00 23.31
                               74.543 38.273 11.543 1.00 28.07
             850 C ARG 266
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                                74.786 39.479 11.517 1.00 29.67
     ATOM
             851 O ARG 266
             852 N TYR 267
                               75.367 37.366 12.053 1.00 28.90
     ATOM
            853 CA TYR 267
                                76.679 37.714 12.558 1.00 30.23
     ATOM
                                77.223 36.584 13.434 1.00 29.98
     ATOM
            854 CB TYR 267
     ATOM
            855 CG TYR 267
                                78.699 36.702 13.727 1.00 31.75
            856 CD1 TYR 267
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                                79.179 37.577 14.712 1.00 31.21
     ATOM
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	ATOM	857 CE1 TYR 267	80.544 37.705 14.950 1.00 31.29
	ATOM	858 CD2 TYR 267	79.625 35.958 12.994 1.00 31.84
	ATOM	859 CE2 TYR 267	80.986 36.078 13.222 1.00 32.15
	ATOM	860 CZ TYR 267	81.442 36.949 14.197 1.00 32.60
5	ATOM	861 OH TYR 267	82.801 37.052 14.389 1.00 34.13
-	ATOM	862 C TYR 267	77.570 37.900 11.343 1.00 31.17
	ATOM	863 O TYR 267	77.543 37.086 10.426 1.00 30.91
	ATOM	864 N ASP 268	78.361 38.966 11.336 1.00 33.09
	ATOM	865 CA ASP 268	79.252 39.233 10.216 1.00 35.57
10	ATOM	866 CB ASP 268	79.085 40.679 9.747 1.00 39.39
	ATOM	867 CG ASP 268	79.796 40.954 8.432 1.00 42.22
	ATOM	868 OD1 ASP 268	79.426 40.331 7.412 1.00 46.07
	ATOM	869 OD2 ASP 268	80.718 41.798 8.415 1.00 44.30
	ATOM	870 C ASP 268	80.700 38.967 10.620 1.00 35.72
15	ATOM	871 O ASP 268	81.287 39.737 11.384 1.00 34.49
	ATOM	872 N PRO 269	81.295 37.872 10.108 1.00 37.00
	ATOM	873 CD PRO 269	80.712 36.887 9.182 1.00 36.77
	ATOM	874 CA PRO 269	82.679 37.514 10.427 1.00 38.52
	ATOM	875 CB PRO 269	82.905 36.239 9.611 1.00 37.06
20	ATOM	876 CG PRO 269	81.549 35.669 9.453 1.00 36.19
	ATOM	877 C PRO 269	83.656 38.613 10.019 1.00 40.96
	ATOM	878 O PRO 269	84.586 38.929 10.760 1.00 42.23
	ATOM	879 N ALA 270	83.418 39.209 8.854 1.00 41.92
	ATOM	880 CA ALA 270	84.277 40.272 8.342 1.00 42.08
25	ATOM	881 CB ALA 270	83.709 40.838 7.029 1.00 42.64
	<b>ATOM</b>	882 C ALA 270	84.495 41.394 9.355 1.00 41.70
	<b>ATOM</b>	883 O ALA 270	85.632 41.709 9.684 1.00 42.25
	ATOM	884 N SER 271	83.408 41.970 9.865 1.00 41.87
	ATOM	885 CA SER 271	83.495 43.073 10.830 1.00 40.75
30	ATOM	886 CB SER 271	82.454 44.143 10.500 1.00 40.60
	ATOM	887 OG SER 271	81.150 43.590 10.464 1.00 40.31
	ATOM	888 C SER 271	83.344 42.658 12.290 1.00 39.99
	ATOM	889 O SER 271	83.484 43.487 13.194 1.00 38.77
	<b>ATOM</b>	890 N ASP 272	83.042 41.381 12.508 1.00 38.94
35	ATOM	891 CA ASP 272	82.859 40.844 13.845 1.00 37.78
	ATOM	892 CB ASP 272	84.182 40.904 14.625 1.00 38.86
	ATOM	893 CG ASP 272	84.094 40.255 16.000 1.00 41.09
	ATOM	894 OD1 ASP 272	83.342 39.275 16.173 1.00 41.64
	ATOM	895 OD2 ASP 272	84.781 40.734 16.924 1.00 43.84
40	ATOM	896 C ASP 272	81.744 41.634 14.536 1.00 36.92
	ATOM	897 O ASP 272	81.907 42.156 15.648 1.00 37.56
	ATOM	898 N THR 273	80.603 41.723 13.865 1.00 33.65
	ATOM	899 CA THR 273	79.469 42.443 14.425 1.00 31.57
•	ATOM	900 CB THR 273	79.246 43.790 13.695 1.00 31.69
45	ATOM	901 OG1 THR 273	79.087 43.557 12.289 1.00 30.71
	ATOM	902 CG2 THR 273	80.426 44.730 13.922 1.00 31.53
	ATOM	903 C THR 273	78.184 41.631 14.310 1.00 30.15
	ATOM	904 O THR 273	78.104 40.697 13.504 1.00 30.10
	ATOM	905 N LEU 274	77.213 41.942 15.164 1.00 27.09
50	ATOM	906 CA LEU 274	75.907 41.303 15.103 1.00 25.94

	ATOM	907 CB LEU 274	75.396 40.936 16.496 1.00 24.47
	ATOM	908 CG LEU 274	76.020 39.731 17.206 1.00 23.33
	ATOM	909 CD1 LEU 274	75.436 39.631 18.602 1.00 21.14
	ATOM	910 CD2 LEU 274	75.792 38.444 16.427 1.00 20.04
5	ATOM	911 C LEU 274	75.010 42.377 14.500 1.00 26.57
	ATOM	912 O LEU 274	75.339 43.557 14.568 1.00 27.03
	ATOM	913 N THR 275	73.914 41.987 13.865 1.00 26.60
	ATOM	914 CA THR 275	73.009 42.966 13.285 1.00 26.48
	<b>ATOM</b>	915 CB THR 275	72.786 42.717 11.781 1.00 26.52
10	<b>ATOM</b>	916 OG1 THR 275	74.044 42.719 11.097 1.00 28.67
	ATOM	917 CG2 THR 275	71.919 43.799 11.198 1.00 27.35
	ATOM	918 C THR 275	71.674 42.898 14.014 1.00 26.57
	ATOM	919 O THR 275	71.069 41.825 14.121 1.00 28.50
	ATOM	920 N LEU 276	71.236 44.026 14.564 1.00 25.18
15	ATOM	921 CA LEU 276	69.970 44.069 15.276 1.00 24.61
	ATOM	922 CB LEU 276	70.057 44.987 16.506 1.00 23.61
	ATOM	923 CG LEU 276	71.199 44.730 17.503 1.00 24.36
	ATOM	924 CD1 LEU 276	71.039 45.654 18.709 1.00 19.91
	ATOM	925 CD2 LEU 276	71.225 43.253 17.947 1.00 22.20
20	ATOM	926 C LEU 276	68.894 44.560 14.322 1.00 25.63
	ATOM	927 O LEU 276	69.100 45.556 13.623 1.00 25.35
	ATOM	928 N SER 277	67.787 43.814 14.249 1.00 25.94
	ATOM	929 CA SER 277	66.634 44.141 13.403 1.00 24.61
	ATOM	930 CB SER 277	65.874 45.335 13.987 1.00 21.96
25	ATOM	931 OG SER 277	65.368 45.029 15.273 1.00 19.68
	ATOM	932 C SER 277	67.005 44.406 11.946 1.00 25.20
	ATOM	933 O SER 277	66.350 45.199 11.267 1.00 25.21
	ATOM	934 N GLY 278 935 CA GLY 278	68.067 43.747 11.489 1.00 27.08 68.556 43.899 10.127 1.00 29.27
30	ATOM ATOM	935 CA GLY 278 936 C GLY 278	69.022 45.297 9.753 1.00 31.57
30	ATOM	937 O GLY 278	69.303 45.564 8.591 1.00 31.42
	ATOM	937 O GL1 278 938 N GLU 279	69.159 46.177 10.740 1.00 33.41
	ATOM	939 CA GLU 279	69.558 47.560 10.484 1.00 34.84
	ATOM	940 CB GLU 279	68.345 48.485 10.650 1.00 36.16
35	ATOM	941 CG GLU 279	67.843 48.606 12.090 1.00 38.08
33	ATOM	942 CD GLU 279	66.566 49.419 12.206 1.00 41.07
	ATOM	943 OE1 GLU 279	66.475 50.279 13.108 1.00 41.98
	ATOM	944 OE2 GLU 279	65.643 49.197 11.399 1.00 43.80
	ATOM	945 C GLU 279	70.706 48.116 11.326 1.00 34.38
40	ATOM	946 O GLU 279	71.366 49.057 10.901 1.00 35.60
. •	ATOM	947 N MET 280	70.944 47.565 12.511 1.00 33.43
	ATOM	948 CA MET 280	72.014 48.085 13.358 1.00 32.27
	ATOM	949 CB MET 280	71.443 48.544 14.702 1.00 31.81
	ATOM	950 CG MET 280	72.471 49.181 15.637 1.00 29.76
45	ATOM	951 SD MET 280	71.813 49.482 17.289 1.00 29.63
	ATOM	952 CE MET 280	70.592 50.735 16.989 1.00 24.91
	<b>ATOM</b>	953 C MET 280	73.161 47.119 13.603 1.00 32.51
	ATOM	954 O MET 280	72.995 46.117 14.303 1.00 32.78
	ATOM	955 N ALA 281	74.321 47.408 13.021 1.00 31.74
50	ATOM	956 CA ALA 281	75.491 46.564 13.231 1.00 32.25

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76.494 46.740 12.108 1.00 30.91
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            958 C ALA 281
     ATOM
    ATOM
            959 O ALA 281
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            961 CA VAL 282
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            962 CB VAL 282
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            963 CG1 VAL 282
            964 CG2 VAL 282
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            966 O VAL 282
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            968 CA LYS 283
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            969 CB LYS 283
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    ATOM
            970 CG LYS 283
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            971 CD LYS 283
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    ATOM
            973 NZ LYS 283
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            974 C LYS 283
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    ATOM
            979 CG ARG 284
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            980 CD ARG 284
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    ATOM
            982 CZ ARG 284
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            984 NH2 ARG 284
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            989 CB GLU 285
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            991 CD GLU 285
                                83.587 49.284 23.747 1.00 54.22
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    ATOM
            992 OE1 GLU 285
                                84.784 49.657 23.760 1.00 55.37
    ATOM
    ATOM
            993 OE2 GLU 285
                                82.686 49.942 23.176 1.00 56.95
            994 C GLU 285
    ATOM
                               80,552 46,785 23.684 1.00 34.45
                               79.990 47.007 24.754 1.00 34.47
    ATOM
            995 O GLU 285
                               80.046 47.166 22.515 1.00 32.27
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    ATOM
            996 N GLN 286
    ATOM
            997 CA GLN 286
                               78,853 47,991 22,438 1.00 30,35
            998 CB GLN 286
                                78.615 48.472 21.006 1.00 33.34
    ATOM
                                79.632 49.497 20.500 1.00 35.09
    ATOM
            999 CG GLN 286
    ATOM 1000 CD GLN 286
                                79.293 50.023 19.108 1.00 38.42
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    ATOM 1001 OE1 GLN 286
                                79.161 49.248 18.158 1.00 39.03
    ATOM 1002 NE2 GLN 286
                                79.156 51.339 18.982 1.00 37.82
                               77.605 47.308 22.970 1.00 29.57
    ATOM 1003 C GLN 286
    ATOM 1004 O GLN 286
                               76.870 47.891 23.770 1.00 26.96
    ATOM 1005 N LEU 287
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                                76.164 45.350 22.979 1.00 28.93
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    ATOM 1006 CA LEU 287
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PCT/US98/25296

### WO 99/26966

	ATOM	1007 CB LEU 287	75.831 44.182 22.029 1.00 27.14
	ATOM	1008 CG LEU 287	74.474 43.484 22.227 1.00 24.66
	ATOM	1009 CD1 LEU 287	73.316 44.475 22.184 1.00 22.70
	ATOM	1010 CD2 LEU 287	74.297 42.413 21.163 1.00 25.17
5	ATOM	1011 C LEU 287	76.303 44.874 24.433 1.00 28.10
	ATOM	1012 O LEU 287	75.301 44.748 25.155 1.00 28.58
	ATOM	1013 N LYS 288	77.541 44.652 24.868 1.00 27.97
	ATOM	1014 CA LYS 288	77.808 44.218 26.230 1.00 28.55
	ATOM	1015 CB LYS 288	79.270 43.800 26.376 1.00 28.93
10	ATOM	1016 CG LYS 288	79.603 43.254 27.750 1.00 32.46
	ATOM	1017 CD LYS 288	81.015 42.725 27.826 1.00 33.48
	ATOM	1018 CE LYS 288	81,205 41.878 29.071 1.00 35.76
	ATOM	1019 NZ LYS 288	82.525 41.186 29.029 1.00 40.52
	ATOM	1020 C LYS 288	77.497 45.341 27.220 1.00 29.15
15	ATOM	1020 C LTS 288	76.782 45.132 28.207 1.00 31.28
15	ATOM	1021 O E15 280 1022 N ASN 289	77.996 46.539 26.933 1.00 28.58
	ATOM	1022 IV ASN 289	77.794 47.692 27.811 1.00 28.40
	ATOM	1025 CA ASN 289	78.815 48.775 27.485 1.00 28.28
	ATOM	1024 CB ASN 289	80.224 48.329 27.770 1.00 31.30
20	ATOM	1025 CO ASIV 289	80.445 47.442 28.601 1.00 33.02
20	ATOM	1027 ND2 ASN 289	81.190 48.928 27.087 1.00 30.49
	ATOM	1027 ND2 ASN 289	76.395 48.278 27.792 1.00 28.33
	ATOM	1028 C ASN 289	76.005 48.977 28.724 1.00 28.36
	ATOM	1030 N GLY 290	75.638 47.977 26.740 1.00 26.71
25	ATOM	1030 N GL1 290	74.286 48.487 26.606 1.00 23.27
23	ATOM	1031 CA GL1 290	73.233 47.852 27.484 1.00 22.93
	ATOM	1032 C GLY 290	72.063 48.219 27.399 1.00 23.84
	ATOM	1033 O GLY 290	73.620 46.905 28.330 1.00 21.30
	ATOM	1034 N GET 291	72.637 46.290 29.199 1.00 20.38
30	ATOM	1035 CA GET 251	72.653 44.778 29.200 1.00 20.05
50	ATOM	1030 C GLY 291	72.190 44.165 30.147 1.00 21.91
	ATOM	1037 O GET 291 1038 N LEU 292	73.211 44.173 28.160 1.00 21.36
	ATOM	1039 CA LEU 292	73.248 42.717 28.062 1.00 21.50
	ATOM	1040 CB LEU 292	73.319 42.280 26.593 1.00 18.52
35	ATOM	1041 CG LEU 292	72.019 42.506 25.815 1.00 17.07
33	ATOM	1041 CO LEO 292 1042 CD1 LEU 292	72.103 41.818 24.479 1.00 18.09
	ATOM	1042 CD1 LEU 292	70.844 41.947 26.599 1.00 16.35
	ATOM	1044 C LEU 292	74.347 42.046 28.872 1.00 22.17
	ATOM	1044 C LEU 292	74.176 40.923 29.352 1.00 21.91
40		1046 N GLY 293	75.479 42.724 29.011 1.00 23.76
40	ATOM ATOM	1046 N GL1 293	76.588 42.169 29.760 1.00 23.70
	ATOM	1047 CA GL1 293	77.134 40.926 29.091 1.00 25.09
			77.362 40.919 27.883 1.00 26.51
	ATOM	1049 O GLY 293 1050 N VAL 294	77.332 39.866 29.867 1.00 26.08
45	ATOM		
45	ATOM	1051 CA VAL 294	77.854 38.618 29.329 1.00 26.34 78.263 37.636 30.443 1.00 26.97
	ATOM	1052 CB VAL 294	79,440 38.199 31.209 1.00 28.20
	ATOM	1053 CG1 VAL 294	77.099 37.371 31.384 1.00 25.56
	ATOM	1054 CG2 VAL 294	76.891 37.937 28.360 1.00 26.41
50	ATOM	1055 C VAL 294 1056 O VAL 294	77.315 37.097 27.568 1.00 27.65
50	ATOM	1056 O VAL 294	11.00 21.00 1.00 21.00

PCT/US98/25296

### WO 99/26966

	ATOM	1057 N VAL 295	75.608 38.304 28.408 1.00 26.09
	<b>ATOM</b>	1058 CA VAL 295	74.606 37.740 27.499 1.00 26.65
	<b>ATOM</b>	1059 CB VAL 295	73.186 38.312 27.777 1.00 28.39
	<b>ATOM</b>	1060 CG1 VAL 295	72.164 37.740 26.782 1.00 26.69
5	<b>ATOM</b>	1061 CG2 VAL 295	72.763 38.005 29.206 1.00 26.23
	<b>ATOM</b>	1062 C VAL 295	75.035 38.089 26.069 1.00 25.83
	<b>ATOM</b>	1063 O VAL 295	74.903 37.286 25.151 1.00 27.12
	<b>ATOM</b>	1064 N SER 296	75.609 39.275 25.908 1.00 24.95
	<b>ATOM</b>	1065 CA SER 296	76.097 39.725 24.619 1.00 26.17
10	ATOM	1066 CB SER 296	76.665 41.132 24.742 1.00 25.82
	<b>ATOM</b>	1067 OG SER 296	77.253 41.554 23.525 1.00 26.64
	<b>ATOM</b>	1068 C SER 296	77.196 38.783 24.142 1.00 28.63
	<b>ATOM</b>	1069 O SER 296	77.241 38.420 22.963 1.00 29.19
	<b>ATOM</b>	1070 N ASP 297	78.118 38.443 25.046 1.00 29.69
15	ATOM	1071 CA ASP 297	79.211 37.531 24.731 1.00 28.96
	ATOM	1072 CB ASP 297	80.058 37.234 25.973 1.00 31.82
	ATOM	1073 CG ASP 297	80.768 38.454 26.506 1.00 35.23
	ATOM	1074 OD1 ASP 297	80.958 39.429 25.743 1.00 35.71
	ATOM	1075 OD2 ASP 297	81.140 38.430 27.698 1.00 37.68
20	ATOM	1076 C ASP 297	78.605 36.227 24.247 1.00 27.63
	<b>ATOM</b>	1077 O ASP 297	79.048 35.666 23.248 1.00 29.88
	ATOM	1078 N ALA 298	77.581 35.762 24.952 1.00 25.15
	ATOM	1079 CA ALA 298	76.909 34.527 24.592 1.00 24.49
	ATOM	1080 CB ALA 298	75.811 34.224 25.594 1.00 21.91
25	ATOM	1081 C ALA 298	76.343 34.569 23.158 1.00 24.93
	ATOM	1082 O ALA 298	76,589 33.654 22.357 1.00 24.83
	ATOM	1083 N ILE 299	75.632 35.647 22.814 1.00 24.70
	ATOM	1084 CA ILE 299	75.041 35.756 21.480 1.00 22.49
	ATOM	1085 CB ILE 299	74.057 36.950 21.351 1.00 21.96
30	ATOM	1086 CG2 ILE 299	73.338 36.876 20.005 1.00 19.17
	ATOM	1087 CG1 ILE 299	72.994 36.876 22.459 1.00 21.16
	<b>ATOM</b>	1088 CD1 ILE 299	72.363 38.228 22.853 1.00 22.04
	<b>ATOM</b>	1089 C ILE 299	76.127 35.829 20.428 1.00 22.33
	ATOM	1090 O ILE 299	75.995 35.234 19.367 1.00 24.80
35	ATOM	1091 N PHE 300	77.209 36.538 20.724 1.00 21.92
	<b>ATOM</b>	1092 CA PHE 300	78.322 36.641 19.785 1.00 23.08
	<b>ATOM</b>	1093 CB PHE 300	79.385 37.636 20.278 1.00 24.08
	<b>ATOM</b>	1094 CG PHE 300	79.249 39.017 19.686 1.00 24.18
	<b>ATOM</b>	1095 CD1 PHE 300	78.494 39.991 20.325 1.00 22.64
40	<b>ATOM</b>	1096 CD2 PHE 300	79.857 39.331 18.471 1.00 23.76
	<b>ATOM</b>	1097 CE1 PHE 300	78.347 41.253 19.770 1.00 22.38
	<b>ATOM</b>	1098 CE2 PHE 300	79.715 40.596 17.904 1.00 23.21
	<b>ATOM</b>	1099 CZ PHE 300	78.957 41.558 18.554 1.00 22.46
	ATOM	1100 C PHE 300	78.948 35.274 19.561 1.00 23.06
45	ATOM	1101 O PHE 300	79.264 34.913 18.426 1.00 23.97
	ATOM	1102 N GLU 301	79.113 34.506 20.636 1.00 23.75
	<b>ATOM</b>	1103 CA GLU 301	79.694 33.169 20.525 1.00 24.16
	<b>ATOM</b>	1104 CB GLU 301	79.884 32.545 21.902 1.00 23.03
	ATOM	1105 C GLU 301	78.776 32.302 19.672 1.00 23.62
50	ATOM	1106 O GLU 301	79.240 31.591 18.777 1.00 25.11

	ATOM	1107 N LEU 302	77.472 32.394 19.926 1.00 23.12
	ATOM	1108 CA LEU 302	76.495 31.624 19.166 1.00 23.56 75.082 31.865 19.701 1.00 21.75
	ATOM	1109 CB LEU 302	
_	ATOM	1110 CG LEU 302	73.953 31.120 18.979 1.00 22.61 74.084 29.612 19.193 1.00 22.31
5	ATOM	1111 CD1 LEU 302 1112 CD2 LEU 302	72.611 31.604 19.485 1.00 19.27
	ATOM	1112 CD2 LEU 302	76.588 32.011 17.687 1.00 24.41
	ATOM ATOM	1113 C LEU 302	76.670 31.140 16.814 1.00 24.63
	ATOM	1114 O LEO 302 1115 N GLY- 303	76.651 33.316 17.425 1.00 25.69
10	ATOM	1116 CA GLY 303	76.746 33.816 16.062 1.00 25.87
10	ATOM	1117 C GLY 303	77.975 33.288 15.338 1.00 28.63
	ATOM	1117 C GLT 303	77.893 32.895 14.170 1.00 28.30
	ATOM	1119 N LYS 304	79.116 33.279 16.023 1.00 29.53
	ATOM	1120 CA LYS 304	80.360 32.791 15.437 1.00 31.18
15	ATOM	1121 CB LYS 304	81.529 32.931 16.418 1.00 34.79
13	ATOM	1122 CG LYS 304	82.157 34.307 16.506 1.00 40.28
	ATOM	1122 CO LTS 304 1123 CD LYS 304	83.441 34.262 17.332 1.00 44.37
	ATOM	1124 CE LYS 304	83.174 33.814 18.775 1.00 47.63
	ATOM	1125 NZ LYS 304	82.459 34.847 19.592 1.00 48.83
20	ATOM	1126 C LYS 304	80.245 31.328 15.042 1.00 30.87
20	ATOM	1127 O LYS 304	80.632 30.944 13.932 1.00 29.53
	ATOM	1128 N SER 305	79.720 30.518 15.961 1.00 30.46
	ATOM	1129 CA SER 305	79.566 29.086 15.731 1.00 31.09
	ATOM	1130 CB SER 305	79.243 28.370 17.041 1.00 29.83
25	<b>ATOM</b>	1131 OG SER 305	77.990 28.783 17.550 1.00 34.66
	ATOM	1132 C SER 305	78.532 28.732 14.653 1.00 31.06
	ATOM	1133 O SER 305	78.745 27.799 13.872 1.00 31.84
	ATOM	1134 N LEU 306	77.436 29.491 14.594 1.00 29.43
	ATOM	1135 CA LEU 306	76.378 29.258 13.611 1.00 28.39
30	ATOM	1136 CB LEU 306	75.121 30.055 13.962 1.00 26.05
	ATOM	1137 CG LEU 306	74.306 29.573 15.157 1.00 26.33
	ATOM	1138 CD1 LEU 306	73.061 30.430 15.285 1.00 26.22
	ATOM	1139 CD2 LEU 306	73.924 28.110 14.985 1.00 25.86
	ATOM	1140 C LEU 306	76.754 29.529 12.157 1.00 28.66
35	ATOM	1141 O LEU 306	76.116 29.001 11.253 1.00 28.58
	ATOM	1142 N SER 307	77.786 30.338 11.931 1.00 29.72
	ATOM	1143 CA SER 307	78.224 30.667 10.577 1.00 31.19
	ATOM	1144 CB SER 307	79.466 31.556 10.617 1.00 30.15 79.226 32.710 11.396 1.00 35.19
40	ATOM	1145 OG SER 307	
40	ATOM	1146 C SER 307	78.531 29.412 9.777 1.00 32.75 78.110 29.283 8.621 1.00 33.09
	ATOM	1147 O SER 307 1148 N ALA 308	79.248 28.482 10.407 1.00 33.36
	ATOM ATOM	1148 N ALA 308 1149 CA ALA 308	79.626 27.223 9.769 1.00 34.50
	ATOM	1150 CB ALA 308	80.636 26.473 10.637 1.00 33.55
45	ATOM	1150 CB ALA 308	78.417 26.328 9.466 1.00 35.00
47	ATOM	1152 O ALA 308	78.469 25.501 8.550 1.00 37.10
	ATOM	1153 N PHE 309	77.335 26.496 10.226 1.00 32.76
	ATOM	1154 CA PHE 309	76.134 25.698 10.028 1.00 31.73
	ATOM	1155 CB PHE 309	75.214 25.818 11.232 1.00 30.04
50	ATOM	1156 CG PHE 309	75.705 25.091 12.438 1.00 31.19

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74.973 24.048 12.975 1.00 31.61
    ATOM 1157 CD1 PHE 309
                                76.884 25.459 13.054 1.00 31.92
    ATOM 1158 CD2 PHE 309
                                75.400 23.391 14.110 1.00 31.22
    ATOM 1159 CE1 PHE 309
    ATOM 1160 CE2 PHE 309
                               77.320 24.807 14.194 1.00 31.01
                               76.577 23.771 14.720 1.00 30.47
    ATOM 1161 CZ PHE 309
                              75.364 26.050 8.753 1.00 31.53
    ATOM 1162 C PHE 309
    ATOM 1163 O PHE 309
                               74.516 25.269 8.310 1.00 31.28
                               75.661 27.220 8.181 1.00 31.12
    ATOM 1164 N ASN 310
                               75.020 27.711 6.957 1.00 30.34
    ATOM 1165 CA ASN 310
10
    ATOM 1166 CB ASN 310
                               75.636 27.036 5.719 1.00 31.63
                               73.511 27.492 7.003 1.00 29.40
    ATOM 1167 C ASN 310
    ATOM 1168 O ASN 310
                               72.939 26.791 6.156 1.00 29.15
    ATOM 1169 N LEU 311
                               72.875 28.055 8.026 1.00 27.60
    ATOM 1170 CA LEU 311
                               71.435 27.907 8.205 1.00 28.23
                               71.021 28.313 9.621 1.00 27.41
    ATOM 1171 CB LEU 311
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    ATOM 1172 CG LEU 311
                               71.603 27.558 10.822 1.00 26.80
    ATOM 1173 CD1 LEU 311
                                70.949 28.078 12.112 1.00 25.05
                              71.360 26.062 10.662 1.00 24.72
    ATOM 1174 CD2 LEU 311
                               70.628 28.719 7.192 1.00 29.01
    ATOM 1175 C LEU 311
    ATOM 1176 O LEU 311
                               71.040 29.808 6.782 1.00 30.66
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    ATOM 1177 N ASP 312
                              69.503 28.168 6.748 1.00 26.30
                               68.675 28.894 5.817 1.00 25.13
    ATOM 1178 CA ASP 312
                               68.391 28.067 4.539 1.00 23.90
    ATOM 1179 CB ASP 312
                               67.438 26.890 4.754 1.00 21.34
    ATOM 1180 CG ASP 312
                                66.959 26.631 5.868 1.00 22.47
25
    ATOM 1181 OD1 ASP 312
    ATOM 1182 OD2 ASP 312
                                67.154 26.206 3.758 1.00 22.18
    ATOM 1183 C ASP 312
                              67.419 29.379 6.542 1.00 24.49
    ATOM 1184 O ASP 312
                               67.221 29.056 7.725 1.00 24.01
    ATOM 1185 N ASP 313
                               66.587 30.153 5.845 1.00 23.40
    ATOM 1186 CA ASP 313
                               65.363 30.697 6.421 1.00 22.63
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                               64.557 31.486 5.385 1.00 24.99
    ATOM 1187 CB ASP 313
                               65.224 32.799 4.994 1.00 28.02
    ATOM 1188 CG ASP 313
                                66.036 33.334 5.778 1.00 30.34
    ATOM 1189 OD1 ASP 313
    ATOM 1190 OD2 ASP 313
                                64.936 33.306 3.897 1.00 30.41
                              64.480 29.650 7.053 1.00 21.47
    ATOM 1191 C ASP 313
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                              63.853 29.917 8.082 1.00 21.76
    ATOM 1192 O ASP 313
    ATOM 1193 N THR 314
                               64.407 28.474 6.435 1.00 19.16
                                63.580 27.386 6.966 1.00 18.79
    ATOM 1194 CA THR 314
    ATOM 1195 CB THR 314
                               63.398 26.240 5.913 1.00 19.68
    ATOM 1196 OG1 THR 314
                                62.743 26.758 4.747 1.00 20.56
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    ATOM 1197 CG2 THR 314
                                62.558 25.112 6.482 1.00 18.84
                               64.133 26.818 8.293 1.00 15.38
    ATOM 1198 C THR 314
    ATOM 1199 O THR 314
                               63.383 26.538 9.223 1.00 14.08
                               65.445 26.656 8.376 1.00 15.16
    ATOM 1200 N GLU 315
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    ATOM 1201 CA GLU 315
                                66.051 26.126 9.593 1.00 16.78
                                67.513 25.785 9.340 1.00 14.29
    ATOM 1202 CB GLU 315
    ATOM 1203 CG GLU 315
                                67.611 24.483 8.579 1.00 15.13
    ATOM 1204 CD GLU 315
                                68.910 24.291 7.872 1.00 15.90
                                69.625 25.285 7.639 1.00 19.80
    ATOM 1205 OE1 GLU 315
                                69.211 23.129 7.527 1.00 19.34
    ATOM 1206 OE2 GLU 315
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65.872 27.119 10.736 1.00 17.27
    ATOM 1207 C GLU 315
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    ATOM 1208 O GLU 315
                               66.081 28.399 10.440 1.00 17.12
    ATOM 1209 N VAL 316
                                65.897 29.441 11.446 1.00 16.92
    ATOM 1210 CA VAL 316
                                66.336 30.828 10.918 1.00 15.89
    ATOM 1211 CB VAL 316
                                66.062 31.921 11.962 1.00 14.60
    ATOM 1212 CG1 VAL 316
                                67.811 30.785 10.579 1.00 15.95
    ATOM 1213 CG2 VAL 316
                               64,430 29,472 11.869 1.00 17.32
    ATOM 1214 C VAL 316
                               64.131 29.582 13.055 1.00 18.11
    ATOM 1215 O VAL 316
                               63.515 29.324 10.905 1.00 17.42
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    ATOM 1216 N ALA 317
                                62.076 29.342 11.195 1.00 16.21
    ATOM 1217 CA ALA 317
                                61.262 29.321 9.910 1.00 14.63
    ATOM 1218 CB ALA 317
                               61.656 28.181 12.079 1.00 16.84
    ATOM 1219 C ALA 317
                               60.904 28.359 13.036 1.00 16.08
    ATOM 1220 O ALA 317
                               62.146 26.990 11.759 1.00 17.27
    ATOM 1221 N LEU 318
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    ATOM 1222 CA LEU 318
                                61.783 25.804 12.526 1.00 17.88
                                62.141 24.525 11.748 1.00 17.58
    ATOM 1223 CB LEU 318
                                61.331 24.333 10.439 1.00 16.87
    ATOM 1224 CG LEU 318
                                61.837 23.155 9.658 1.00 15.79
    ATOM 1225 CD1 LEU 318
                                59.860 24.149 10.728 1.00 14.08
     ATOM 1226 CD2 LEU 318
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                               62.394 25.852 13.932 1.00 18.20
     ATOM 1227 C LEU 318
                               61.733 25.495 14.910 1.00 18.71
    ATOM 1228 O LEU 318
                               63.614 26.380 14.034 1.00 17.73
     ATOM 1229 N LEU 319
                                64.288 26.531 15.321 1.00 16.57
     ATOM 1230 CA LEU 319
                                65.689 27.105 15.107 1.00 18.81
25
     ATOM 1231 CB LEU 319
                                66.733 27.223 16.224 1.00 21.77
     ATOM 1232 CG LEU 319
                                66.767 25.994 17.117 1.00 23.03
     ATOM 1233 CD1 LEU 319
                                68.076 27.421 15.554 1.00 20.86
     ATOM 1234 CD2 LEU 319
                               63.433 27.471 16.160 1.00 16.07
     ATOM 1235 C LEU 319
                               63.134 27.183 17.319 1.00 16.40
     ATOM 1236 O LEU 319
30
                               62.948 28.546 15.545 1.00 13.91
     ATOM 1237 N GLN 320
                                62.101 29.490 16.253 1.00 13.86
     ATOM 1238 CA GLN 320
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     ATOM 1239 CB GLN 320
                                62.994 31.553 15.080 1.00 12.17
     ATOM 1240 CG GLN 320
                                62.691 32.802 14.253 1.00 13.98
     ATOM 1241 CD GLN 320
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                                63.597 33.568 13.950 1.00 15.61
     ATOM 1242 OE1 GLN 320
                                61.436 32.993 13.862 1.00 13.85
     ATOM 1243 NE2 GLN 320
                               60.813 28.832 16.746 1.00 14.52
     ATOM 1244 C GLN 320
                               60.367 29.087 17.864 1.00 15.12
     ATOM 1245 O GLN 320
                               60.211 27.982 15.924 1.00 14.21
     ATOM 1246 N ALA 321
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     ATOM 1247 CA ALA 321
                                58.408 26.519 15.115 1.00 13.84
     ATOM 1248 CB ALA 321
                               59.217 26.349 17.487 1.00 15.98
     ATOM 1249 C ALA 321
                                58.358 26.197 18.355 1.00 15.12
     ATOM 1250 O ALA 321
                               60.373 25.687 17.488 1.00 16.63
     ATOM 1251 N VAL 322
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                                60.720 24.757 18.557 1.00 18.74
     ATOM 1252 CA VAL 322
                                62.012 23.943 18.231 1.00 19.42
     ATOM 1253 CB VAL 322
                                 62.493 23.154 19.455 1.00 19.45
     ATOM 1254 CG1 VAL 322
     ATOM 1255 CG2 VAL 322
                                 61.745 22.986 17.083 1.00 19.05
                               60.910 25.556 19.833 1.00 18.42
     ATOM 1256 C VAL 322
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	4 TO) 4	1257 O VAL 322	60.421 25.164 20.886 1.00 19.46
	ATOM		61.607 26.685 19.735 1.00 18.65
	ATOM		61.836 27.543 20.894 1.00 18.49
	ATOM		62,710 28.740 20.508 1.00 18.36
_	ATOM	1260 CB LEU 323	64.179 28.449 20.186 1.00 18.13
5	ATOM	1261 CG LEU 323	64.829 29.669 19.585 1.00 17.37
	ATOM	1262 CD1 LEU 323	64.923 27.999 21.447 1.00 17.27
	ATOM	1263 CD2 LEU 323	
	ATOM	1264 C LEU 323	60.499 28.029 21.454 1.00 18.38
	ATOM	1265 O LEU 323	60.275 28.008 22.663 1.00 18.81
10	ATOM	1266 N LEU 324	59.595 28.406 20.557 1.00 18.67
	ATOM	1267 CA LEU 324	58.275 28.897 20.924 1.00 19.02
	ATOM	1268 CB LEU 324	57.564 29.467 19.685 1.00 17.78
	ATOM	1269 CG LEU 324	56.095 29.891 19.838 1.00 17.59
	ATOM	1270 CD1 LEU 324	55.983 31.123 20.709 1.00 18.15
15	ATOM	1271 CD2 LEU 324	55.489 30.180 18.476 1.00 16.43
	ATOM	1272 C LEU 324	57.354 27.884 21.610 1.00 19.62
	ATOM	1273 O LEU 324	56.735 28.185 22.633 1.00 19.40
	ATOM	1274 N MET 325	57.224 26.701 21.029 1.00 21.14
20	ATOM	1275 CA MET 325	56.330 25.680 21.585 1.00 24.06 55 857 24.738 20.473 1.00 24.68
20	ATOM	1276 CB MET 325	20.007 211720 207170 11110 11110
	ATOM	1277 CG MET 325	55.169 25.444 19.303 1.00 24.49 53.759 26.457 19.820 1.00 26.18
	ATOM	1278 SD MET 325	52.609 25.252 20.373 1.00 24.03
	ATOM	1279 CE MET 325	56.996 24.887 22.705 1.00 26.15
25	ATOM	1280 C MET 325	57.021 23.664 22.693 1.00 25.68
25	ATOM	1281 O MET 325 1282 N SER 326	57.555 25.593 23.671 1.00 29.34
	ATOM ATOM	1282 N SER 326	58.232 24.938 24.774 1.00 32.40
	ATOM	1284 CB SER 326	59.512 25.701 25.112 1.00 32.12
	ATOM	1285 OG SER 326	60.127 25.173 26.272 1.00 36.86
30	ATOM	1286 C SER 326	57.317 24.831 25.996 1.00 34.04
30	ATOM	1287 O SER 326	56.532 25.741 26.280 1.00 33.24
	ATOM	1288 N THR 327	57.366 23.687 26.674 1.00 35.62
	ATOM	1289 CA THR 327	56.560 23.486 27.867 1.00 36.88
	ATOM	1290 CB THR 327	55.938 22.085 27.907 1.00 36.58
35	ATOM	1291 OG1 THR 327	56.953 21.094 27.714 1.00 38.58
55	ATOM	1292 CG2 THR 327	54.883 21.938 26.826 1.00 37.73
	ATOM	1293 C THR 327	57.378 23.733 29.135 1.00 38.77
	ATOM	1294 O THR 327	56.921 23.438 30.240 1.00 39.53
	ATOM	1295 N ASP 328	58.593 24.260 28.972 1.00 41.25
40	ATOM	1296 CA ASP 328	59.473 24.573 30.099 1.00 43.20
	ATOM	1297 CB ASP 328	60.940 24.698 29.655 1.00 46.47
	ATOM	1298 CG ASP 328	61.618 23.346 29.439 1.00 51.94
	ATOM	1299' OD1 ASP 328	62.547 23.278 28.601 1.00 55.43
	ATOM	1300 OD2 ASP 328	61.251 22.354 30.111 1.00 54.77
45	ATOM	1301 C ASP 328	59.001 25.905 30.653 1.00 43.79
1,5	ATOM	1302 O ASP 328	59.755 26.877 30.709 1.00 45.91
	ATOM	1303 N ARG 329	57.724 25.967 30.995 1.00 43.55
	ATOM	1304 CA ARG 329	57.143 27.178 31.542 1.00 43.04
	ATOM	1305 CB ARG 329	56.398 27.997 30.482 1.00 43.87
50	ATOM	1306 CG ARG 329	57,258 28.740 29.504 1.00 40.87
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WO 99/26966

## PCT/US98/25296

	ATOM	1307 CD ARG 329	57.545 27.886 28.314 1.00 39.52
	ATOM	1308 NE ARG 329	58.301 28.643 27.341 1.00 38.90
	ATOM	1309 CZ ARG 329	59.624 28.708 27.313 1.00 40.59
	ATOM	1310 NH1 ARG 329	60.359 28.052 28.196 1.00 42.41
5	ATOM	1311 NH2 ARG 329	60.210 29.466 26.413 1.00 41.87
•	ATOM	1312 C ARG 329	56.152 26.817 32.609 1.00 43.00
	ATOM	1313 O ARG 329	55.600 25.716 32.628 1.00 43.66
	ATOM	1314 N SER 330	55.886 27.797 33.456 1.00 41.58
	ATOM	1315 CA SER 330	54.953 27.641 34.538 1.00 40.11
10	ATOM	1316 CB SER 330	55.491 28.362 35.777 1.00 40.38
	ATOM	1317 C SER 330	53.602 28.223 34.103 1.00 38.99
	ATOM	1318 O SER 330	53.553 29.172 33.320 1.00 39.22
	ATOM	1319 N GLY 331	52.517 27.581 34.529 1.00 37.52
	ATOM	1320 CA GLY 331	51.176 28.063 34.232 1.00 35.64
15	ATOM	1321 C GLY 331	50.493 27.782 32.906 1.00 35.14
15	ATOM	1322 O GLY 331	49.439 28.363 32.640 1.00 34.48
	ATOM	1323 N LEU 332	51.059 26.925 32.066 1.00 34.54
	ATOM	1324 CA LEU 332	50.424 26.637 30.780 1.00 34.59
	ATOM	1325 CB LEU 332	51.394 25.942 29.828 1.00 33.09
20	ATOM	1326 CG LEU 332	52.532 26.765 29.236 1.00 32.72
20	ATOM	1327 CD1 LEU 332	53.473 25.834 28.497 1.00 30.29
	ATOM	1327 CD1 LEU 332	51.987 27.844 28.313 1.00 29.20
	ATOM	1329 C LEU 332	49.191 25.763 30.969 1.00 35.14
	ATOM	1330 O LEU 332	49.178 24.874 31.811 1.00 35.96
25	ATOM	1331 N LEU 333	48.153 26.076 30.204 1.00 35.65
23	ATOM	1332 CA LEU 333	46.898 25.345 30.215 1.00 37.97
	ATOM	1333 CB LEU 333	45.743 26.271 29.796 1.00 40.71
	ATOM	1334 CG LEU 333	45.389 27.483 30.670 1.00 43.46
	ATOM ATOM	1335 CD1 LEU 333	44.713 28.620 29.882 1.00 42.72
30	ATOM	1336 CD2 LEU 333	44.487 27.021 31.806 1.00 45.25
50	ATOM	1337 C LEU 333	46.952 24.115 29.300 1.00 37.78
	ATOM	1338 O LEU 333	46.695 22.991 29.720 1.00 37.65
	ATOM	1339 N CYA 334	47.361 24.323 28.060 1.00 38.65
	ATOM	1340 CA CYA 334	47.413 23.249 27.073 1.00 40.91
35	ATOM	1341 CB CYA 334	46.936 23.788 25.721 1.00 47.35
<i>J J</i>	ATOM	1342 SG CYA 334	45.406 24.693 25.867 1.00 52.24
	ATOM	1343 AS CYA 334	44.066 22.890 25.562 1.00 70.72
	ATOM	1344 C CYA 334	48.778 22.588 26.901 1.00 39.85
	ATOM	1345 O CYA 334	49.287 22.473 25.775 1.00 39.54
40		1346 N VAL 335	49.329 22.078 27.997 1.00 37.67
-10	ATOM	1347 CA VAL 335	50.641 21.432 27.967 1.00 36.07
	ATOM	1348 CB VAL 335	51.019 20.905 29.384 1.00 33.70
	ATOM	1349 CG1 VAL 335	52.434 20.332 29.401 1.00 33.70
	ATOM	1350 CG2 VAL 335	50.913 22.028 30.387 1.00 31.84
45	ATOM	1351 C VAL 335	50.734 20.334 26.885 1.00 36.09
73	ATOM	1352 O VAL 335	51.662 20.335 26.064 1.00 34.41
	ATOM	1352 O VAL 335	49.747 19.444 26.833 1.00 35.95
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	ATOM	1355 CB ASP 336	48.591 17.394 26.091 1.00 41.36
50	ATOM	1356 CG ASP 336	48.613 16.206 25.129 1.00 46.23
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     ATOM 1360 O ASP 336
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     ATOM 1362 CA LYS 337
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     ATOM 1363 CB LYS 337
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     ATOM 1365 O LYS 337
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                              50.597 21.688 23.208 1.00 25.90
     ATOM 1366 N ILE 338
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     ATOM 1368 CB ILE 338
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     ATOM 1370 CG1 ILE 338
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     ATOM 1377 C GLU 339
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     ATOM 1378 O GLU 339
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     ATOM 1381 CB LYS 340
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     ATOM 1382 C LYS 340
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     ATOM 1383 O LYS 340
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     ATOM 1385 CA SER 341
     ATOM 1386 CB SER 341
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     ATOM 1389 O SER 341
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     ATOM 1393 CG GLN 342
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                                59.476 21.677 22.057 1.00 26.93
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     ATOM 1395 OE1 GLN 342
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     ATOM 1397 C GLN 342
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                                57.222 17.280 18.610 1.00 25.34
     ATOM 1400 CA GLU 343
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     ATOM 1404 N ALA 344
     ATOM 1405 CA ALA 344
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     ATOM 1406 CB ALA 344
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	ATOM	1407 C ALA 344	56.768 18.743 14.489 1.00 22.77
	ATOM	1408 O ALA 344	57.355 18.360 13.477 1.00 22.08
	ATOM	1409 N TYR 345	57.057 19.893 15.092 1.00 21.89
	ATOM	1410 CA TYR 345	58.075 20.792 14.550 1.00 21.18
5	ATOM	1411 CB TYR 345	58.108 22.119 15.313 1.00 20.27
_	ATOM	1412 CG TYR 345	57.048 23.078 14.856 1.00 17.45
	ATOM	1413 CD1 TYR 345	56,001 23.431 15.698 1.00 17.99
	ATOM	1414 CE1 TYR 345	54.992 24.253 15.270 1.00 19.97
	ATOM	1415 CD2 TYR 345	57.063 23.589 13.562 1.00 19.11
10	ATOM	1416 CE2 TYR 345	56.055 24.424 13.116 1.00 19.14
10	ATOM	1417 CZ TYR 345	55.017 24.749 13.972 1.00 20.78
	ATOM	1418 OH TYR 345	53.983 25.539 13.530 1.00 20.70
	ATOM	1419 C TYR 345	59.454 20.167 14.583 1.00 20.96
	ATOM	1420 O TYR 345	60.221 20.314 13.632 1.00 22.29
15	ATOM	1421 N LEU 346	59.778 19.480 15.677 1.00 20.82
15	ATOM	1422 CA LEU 346	61.079 18.838 15.817 1.00 20.18
	ATOM	1423 CB LEU 346	61.216 18.203 17.205 1.00 21.04
	ATOM	1424 CG LEU 346	61.606 19.158 18.335 1.00 21.25
	ATOM	1425 CD1 LEU 346	61.226 18.595 19.685 1.00 20.95
20	ATOM	1426 CD2 LEU 346	63.099 19.438 18.267 1.00 19.90
20	ATOM	1427 C LEU 346	61.317 17.806 14.716 1.00 20.19
	ATOM	1428 O LEU 346	62.407 17.755 14.142 1.00 20.69
	ATOM	1429 N LEU 347	60.290 17.016 14.390 1.00 22.00
	ATOM	1430 CA LEU 347	60.406 15.994 13.344 1.00 21.81
25	ATOM	1431 CB LEU 347	59.199 15.051 13.366 1.00 24.03
	ATOM	1432 CG LEU 347	59.301 13.805 14.250 1.00 26.28
	ATOM	1433 CD1 LEU 347	57.964 13.072 14.277 1.00 27.79
	ATOM	1434 CD2 LEU 347	60.409 12.889 13.728 1.00 24.78
	ATOM	1435 C LEU 347	60.544 16.623 11.966 1.00 20.50
30	ATOM	1436 O LEU 347	61.351 16.179 11.143 1.00 21.39
	ATOM	1437 N ALA 348	59.767 17.674 11.727 1.00 20.84
	<b>ATOM</b>	1438 CA ALA 348	59.788 18.381 10.456 1.00 18.12
	ATOM	1439 CB ALA 348	58.729 19.480 10.457 1.00 18.49
	<b>ATOM</b>	1440 C ALA 348	61.168 18.963 10.269 1.00 17.53
35	<b>ATOM</b>	1441 O ALA 348	61.785 18.781 9.228 1.00 18.78
	<b>ATOM</b>	1442 N PHE 349	61.677 19.569 11.338 1.00 19.55
	ATOM	1443 CA PHE 349	63.001 20.196 11.389 1.00 19.84
	ATOM	1444 CB PHE 349	63.188 20.823 12.786 1.00 18.68
	ATOM	1445 CG PHE 349	64.380 21.758 12.917 1.00 19.12
40	ATOM	1446 CD1 PHE 349	65.234 22.008 11.851 1.00 19.95
	ATOM	1447 CD2 PHE 349	64.618 22.420 14.126 1.00 20.06
	ATOM	1448 CE1 PHE 349	66.294 22.905 11.971 1.00 18.99
	ATOM	1449 CE2 PHE 349	65.674 23.317 14.261 1.00 16.79
	ATOM	1450 CZ PHE 349	66.516 23.562 13.184 1.00 18.91
45	ATOM	1451 C PHE 349	64.108 19.170 11.103 1.00 20.44
	ATOM	1452 O PHE 349	64.980 19.401 10.260 1.00 19.83
	ATOM	1453 N GLU 350	64.064 18.032 11.794 1.00 23.59
	ATOM	1454 CA GLU 350	65.077 16.995 11.610 1.00 23.46
	ATOM	1455 CB GLU 350	64.830 15.845 12.584 1.00 25.26
50	ATOM	1456 CG GLU 350	65.694 14.644 12.288 1.00 31.98

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ATOM 1457 CD GLU 350
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     ATOM 1458 OE1 GLU 350
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     ATOM 1459 OE2 GLU 350
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     ATOM 1460 C GLU 350
                               65.083 16.489 10.165 1.00 21.12
     ATOM 1461 O GLU 350
                               66.133 16.384 9.526 1.00 19.81
                               63.888 16.234 9.651 1.00 21.98
     ATOM 1462 N HIS 351
                               63.694 15.751 8.292 1.00 21.31
     ATOM 1463 CA HIS 351
     ATOM 1464 CB HIS 351
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     ATOM 1465 CG HIS 351
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     ATOM 1466 CD2 HIS 351
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     ATOM 1467 ND1 HIS 351
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     ATOM 1470 C HIS 351
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                                65.132 22.507 6.438 1.00 18.91
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     ATOM 1479 CE2 TYR 352
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     ATOM 1485 CA VAL 353
    ATOM 1486 CB VAL 353
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    ATOM 1488 CG2 VAL 353
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    ATOM 1493 CB ASN 354
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    ATOM 1494 CG ASN 354
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    ATOM 1503 CD2 HIS 355
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    ATOM 1506 NE2 HIS 355
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PCT/US98/25296

### WO 99/26966

	ATOM	1507 C HIS 355	68.022 17.857 2.624 1.00 24.07
	ATOM	1508 O HIS 355	68.460 17.729 1.484 1.00 23.54
	ATOM	1509 N ARG 356	68.463 18.774 3.471 1.00 25.31
	ATOM	1510 CA ARG 356	69.523 19.714 3.130 1.00 25.69
5	ATOM	1511 CB ARG 356	69.561 20.820 4.168 1.00 24.06
	ATOM	1512 CG ARG 356	68.337 21.682 4.094 1.00 23.23
	ATOM	1513 CD ARG 356	68.670 22.973 3.424 1.00 25.91
	ATOM	1514 NE ARG 356	69.447 23.814 4.322 1.00 24.87
	<b>ATOM</b>	1515 CZ ARG 356	70.325 24.726 3.928 1.00 25.05
10	ATOM	1516 NH1 ARG 356	70,546 24,920 2.640 1.00 24.97
	ATOM	1517 NH2 ARG 356	
	ATOM	1518 C ARG 356	70.900 19.109 2.949 1.00 27.73
	ATOM	1519 O ARG 356	71.724 19.645 2.208 1.00 28.38
	ATOM	1520 N LYS 357	71.179 18.048 3.693 1.00 29.45
15	ATOM	1521 CA LYS 357	72.457 17.355 3.588 1.00 31.35
13	ATOM	1522 CB LYS 357	72.503 16.566 2.270 1.00 32.80
	ATOM	1523 CG LYS 357	71.290 15.650 2.103 1.00 35.78
	ATOM	1524 CD LYS 357	71.264 14.927 0.778 1.00 39.43
	ATOM	1525 CE LYS 357	70.121 13.918 0.739 1.00 42.93
20	ATOM	1526 NZ LYS 357	70.162 13.074 -0.498 1.00 45.97
20	ATOM	1527 C LYS 357	73.692 18.247 3.743 1.00 31.34
	ATOM	1528 O LYS 357	74.489 18.390 2.818 1.00 32.65
	ATOM	1529 N HIS 358	73.837 18.861 4.913 1.00 30.72
	<del>-</del>	1530 CA HIS 358	74.995 19.706 5.186 1.00 31.49
25	ATOM	1531 CB HIS 358	74.895 20.322 6.579 1.00 29.13
25	ATOM	1531 CB HIS 358	73.882 21.415 6.688 1.00 25.30
	ATOM	1532 CO HIS 358 1533 CD2 HIS 358	74.026 22.760 6.646 1.00 24.90
	ATOM	1534 ND1 HIS 358	72.543 21.175 6.892 1.00 24.54
	ATOM		71.901 22.324 6.975 1.00 24.34
20	ATOM		72.777 23.302 6.830 1.00 25.28
30	ATOM	1536 NE2 HIS 358	76.235 18.831 5.161 1.00 33.38
	ATOM	1537 C HIS 358	
	ATOM	1538 O HIS 358	76.166 17.647 5.495 1.00 35.46
	ATOM	1539 N ASN 359	77.366 19.399 4.768 1.00 35.34
	ATOM	1540 CA ASN 359	78.606 18.636 4.746 1.00 38.17
35	ATOM	1541 CB ASN 359	79.544 19.150 3.646 1.00 37.84
	ATOM	1542 C ASN 359	79.236 18.825 6.120 1.00 39.85
	ATOM	1543 O ASN 359	80.317 19.406 6.240 1.00 42.72
	ATOM	1544 N ILE 360	78.510 18.411 7.159 1.00 39.01
	ATOM	1545 CA ILE 360	78.968 18.526 8.549 1.00 36.72
40	ATOM	1546 CB ILE 360	78.351 19.752 9.264 1.00 37.69
	ATOM	1547 CG2 ILE 360	78.802 19.793 10.722 1.00 37.56
	ATOM	1548 CG1 ILE 360	78.735 21.049 8.549 1.00 37.68
	ATOM	1549 CD1 ILE 360	77.970 22.253 9.041 1.00 38.40
	ATOM	1550 C ILE 360	78.524 17.278 9.303 1.00 35.15
45	ATOM	1551 O ILE 360	77.343 16.931 9.314 1.00 33.75
	ATOM	1552 N PRO 361	79.475 16.564 9.912 1.00 34.64
	ATOM	1553 CD PRO 361	80.930 16.785 9.873 1.00 35.59
	ATOM	1554 CA PRO 361	79.138 15.349 10.660 1.00 33.92
	ATOM	1555 CB PRO 361	80.513 14.768 11.014 1.00 35.27
50	ATOM	1556 CG PRO 361	81.412 15.972 11.048 1.00 35.97

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     ATOM 1562 CG HIS 362
                                78.075 13.441 14.440 1.00 33.72
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                               75.521 16.769 14.472 1.00 29.93
     ATOM 1568 O HIS 362
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     ATOM 1582 CB TRP 364
                                70.444 16.426 13.506 1.00 24.27
     ATOM 1583 CG TRP 364
                                69.168 16.391 14.328 1.00 23.75
     ATOM 1584 CD2 TRP 364
                                68.152 17.407 14.397 1.00 24.87
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     ATOM 1586 CE3 TRP 364
                                67.989 18.674 13.820 1.00 25.47
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     ATOM 1588 NE1 TRP 364
                                67.530 15.679 15.684 1.00 25.99
     ATOM 1589 CZ2 TRP 364
                                65.987 17.661 15.560 1.00 25.14
     ATOM 1590 CZ3 TRP 364
                                66.844 19.405 14.116 1.00 25.29
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     ATOM 1591 CH2 TRP 364
                                65.857 18.894 14.982 1.00 24.53
     ATOM 1592 C TRP 364
                               71.659 17.747 15.308 1.00 26.94
     ATOM 1593 O TRP 364
                               71.202 18.721 15.904 1.00 27.16
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                               72.382 16.796 15.944 1.00 27.60
                                72.912 15.522 15.411 1.00 27.55
     ATOM 1595 CD PRO 365
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     ATOM 1596 CA PRO 365
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                                73.565 15.717 17.668 1.00 26.00
    ATOM 1598 CG PRO 365
                                73.136 14.705 16.658 1.00 28.32
    ATOM 1599 C PRO 365
                               73.374 18.225 17.714 1.00 23.89
     ATOM 1600 O PRO 365
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                               74,297 18.626 16.845 1.00 24.24
    ATOM 1602 CA LYS 366
                                75.058 19.862 17.027 1.00 26.24
    ATOM 1603 CB LYS 366
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     ATOM 1604 CG LYS 366
                                77.310 19.022 16.138 1.00 28.76
     ATOM 1605 CD LYS 366
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    ATOM 1606 CE LYS 366
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	ATOM	1607 NZ LYS 366	80.388 18.463 13.947 1.00 37.89
	ATOM	1608 C LYS 366	74.181 21.107 16.993 1.00 26.73
	ATOM	1609 O LYS 366	74.385 22.042 17.762 1.00 27.36
	ATOM	1610 N LEU 367	73.216 21.124 16.086 1.00 27.98
5	ATOM	1611 CA LEU 367	72.308 22.256 15.967 1.00 27.87
•	ATOM	1612 CB LEU 367	71.559 22.192 14.632 1.00 27.29
	ATOM	1613 CG LEU 367	70.613 23.356 14.318 1.00 27.25
	ATOM	1614 CD1 LEU 367	71.334 24.707 14.510 1.00 22.90
	ATOM	1615 CD2 LEU 367	70.081 23.189 12.896 1.00 24.54
10	ATOM	1616 C LEU 367	71.327 22.223 17.134 1.00 29.38
••	ATOM	1617 O LEU 367	70.993 23.249 17.716 1.00 31.09
	ATOM	1618 N LEU 368	70.889 21.026 17.491 1.00 30.38
	ATOM	1619 CA LEU 368	69.962 20.843 18.594 1.00 31.14
	ATOM	1620 CB LEU 368	69.659 19.353 18.731 1.00 32.20
15	ATOM	1621 CG LEU 368	68.247 18.852 19.014 1.00 33.52
13	ATOM	1622 CD1 LEU 368	67.184 19.651 18.267 1.00 31.14
	ATOM	1623 CD2 LEU 368	68.210 17.379 18.632 1.00 33.99
	ATOM	1624 C LEU 368	70.601 21.395 19.876 1.00 32.36
	ATOM	1625 O LEU 368	69.917 21.963 20.730 1.00 32.58
20	ATOM	1626 N MET 369	71.922 21.272 19.985 1.00 33.30
20	ATOM	1627 CA MET 369	72.641 21.771 21.149 1.00 34.04
	ATOM	1628 CB MET 369	74.051 21.190 21.209 1.00 35.31
	ATOM	1629 CG MET 369	74.108 19.858 21.935 1.00 36.83
	ATOM	1630 SD MET 369	75.312 18.728 21.235 1.00 43.07
25	ATOM	1631 CE MET 369	76.862 19.636 21.472 1.00 41.31
	ATOM	1632 C MET 369	72.675 23.297 21.212 1.00 34.30
	ATOM	1633 O MET 369	72.961 23.876 22.269 1.00 35.82
	ATOM	1634 N LYS 370	72.368 23.949 20.091 1.00 32.14
	ATOM	1635 CA LYS 370	72.325 25.405 20.044 1.00 29.17
30	ATOM	1636 CB LYS 370	72.394 25.904 18.608 1.00 28.18
	ATOM	1637 CG LYS 370	73.662 25.518 17.900 1.00 27.72
	ATOM	1638 CD LYS 370	74.866 25.969 18.679 1.00 28.10
	ATOM	1639 CE LYS 370	76.127 25.650 17.930 1.00 27.79
	ATOM	1640 NZ LYS 370	77.298 25.941 18.777 1.00 30.78
35	ATOM	1641 C LYS 370	71.033 25.875 20.705 1.00 29.27
	ATOM	1642 O LYS 370	70.950 26.999 21.200 1.00 29.43
	ATOM	1643 N VAL 371	70.018 25.014 20.714 1.00 29.40
	ATOM	1644 CA VAL 371	68.756 25.358 21.358 1.00 29.90
	ATOM	1645 CB VAL 371	67.687 24.237 21.218 1.00 28.75
40	ATOM		66.463 24.561 22.064 1.00 27.12
	ATOM	1647 CG2 VAL 371	67.275 24.080 19.762 1.00 29.23
	ATOM	1648 C VAL 371	69.075 25.573 22.832 1.00 31.39
	ATOM	1649 O VAL 371	68.543 26.481 23.462 1.00 31.20
	ATOM	1650 N THR 372	69.971 24.743 23.366 1.00 31.39
45	ATOM	1651 CA THR 372	70.371 24.847 24.762 1.00 31.10
	ATOM	1652 CB THR 372	71,282 23,664 25,170 1.00 31.59
	ATOM	1653 OG1 THR 372	70.554 22.441 25.008 1.00 30.60
	ATOM	1654 CG2 THR 372	71.720 23.795 26.625 1.00 30.14
	ATOM	1655 C THR 372	71.071 26.186 24.994 1.00 30.76
50	ATOM	1656 O THR 372	70.711 26.935 25.910 1.00 31.45

WO 99/26966

PCT/US98/25296

	ATOM	1657 N ASP 373	72.038 26.507 24.138 1.00 29.31
	<b>ATOM</b>	1658 CA ASP 373	72.744 27.772 24.252 1.00 27.32
	ATOM	1659 CB ASP 373	73.745 27.934 23.115 1.00 27.98
	ATOM	1660 CG ASP 373	74.886 26.933 23.190 1.00 28.94
5	ATOM	1661 OD1 ASP 373	75.043 26.259 24.225 1.00 31.01
	ATOM	1662 OD2 ASP 373	75.639 26.825 22.205 1.00 31.38
	ATOM	1663 C ASP 373	71.742 28.926 24.247 1.00 26.50
	ATOM	1664 O ASP 373	71.872 29.861 25.040 1.00 27.35
	ATOM	1665 N LEU 374	70.711 28.826 23.412 1.00 24.17
10	ATOM	1666 CA LEU 374	69.688 29.864 23.331 1.00 23.38
	ATOM	1667 CB LEU 374	68.795 29.660 22.107 1.00 22.98
	ATOM	1668 CG LEU 374	69.361 30.183 20.786 1.00 24.45
	ATOM	1669 CD1 LEU 374	68.668 29.520 19.589 1.00 24.72
	ATOM	1670 CD2 LEU 374	69.223 31.704 20.735 1.00 22.40
15	ATOM	1671 C LEU 374	68.839 29.964 24.589 1.00 24.31
13	ATOM	1672 O LEU 374	68.442 31.065 24.986 1.00 23.31
	ATOM	1673 N ARG 375	68.543 28.826 25.211 1.00 25.32
	ATOM	1674 CA ARG 375	67.748 28.821 26.438 1.00 27.76
	ATOM	1675 CB ARG 375	67.455 27.392 26.908 1.00 30.82
20	ATOM	1676 CG ARG 375	66.901 26.439 25.854 1.00 38.79
20	ATOM	1677 CD ARG 375	65.424 26.630 25.582 1.00 45.40
	ATOM	1678 NE ARG 375	64.709 25.360 25.620 1.00 52.61
	ATOM	1679 CZ ARG 375	63.800 24.967 24.726 1.00 56.89
	ATOM	1680 NH1 ARG 375	63.473 25.732 23.694 1.00 58.27
25	ATOM	1681 NH2 ARG 375	63.201 23.793 24.855 1.00 58.46
	ATOM	1682 C ARG 375	68.563 29.542 27.512 1.00 26.98
	ATOM	1683 O ARG 375	68.025 30.336 28.282 1.00 26.18
	ATOM	1684 N MET 376	69.862 29.255 27.551 1.00 26.80
	ATOM	1685 CA MET 376	70.767 29.867 28.511 1.00 29.22
30	ATOM	1686 CB MET 376	72.172 29.270 28.379 1.00 33.70
	ATOM	1687 CG MET 376	72.595 28.371 29.562 1.00 43.20
	ATOM	1688 SD MET 376	73,320 29.260 31.011 1.00 52.38
	ATOM	1689 CE MET 376	71.843 29.854 31.913 1.00 48.11
	ATOM	1690 C MET 376	70.804 31.384 28.339 1.00 27.54
35	ATOM	1691 O MET 376	70.792 32.126 29.323 1.00 26.96
_	ATOM		70.841 31.835 27.087 1.00 25.39
	ATOM	1693 CA ILE 377	70.847 33.264 26.767 1.00 23.26
	ATOM	1694 CB ILE 377	70.992 33.488 25.222 1.00 22.73
	ATOM	1695 CG2 ILE 377	70.560 34.909 24.819 1.00 21.81
40	<b>ATOM</b>	1696 CG1 ILE 377	72.431 33.205 24.789 1.00 20.39
	ATOM	1697 CD1 ILE 377	72.644 33.148 23.300 1.00 18.85
	ATOM		69.558 33.900 27.309 1.00 22.91
	ATOM		69.597 34.925 27.989 1.00 22.02
	ATOM	1700 N GLY 378	68.427 33.244 27.069 1.00 22.29
45	ATOM	1701 CA GLY 378	67.161 33.757 27.547 1.00 22.83
	ATOM	1702 C GLY 378	67.111 33.815 29.063 1.00 25.60
	ATOM	1703 O GLY 378	66.546 34.752 29.630 1.00 26.25
	ATOM	1704 N ALA 379	67.691 32.804 29.713 1.00 26.88
	ATOM	1705 CA ALA 379	67.744 32.707 31.175 1.00 27.19
50	ATOM	1706 CB ALA 379	68.322 31.358 31.590 1.00 26.97

. WO 99/26966

# PCT/US98/25296

	ATOM	1707 C ALA 379	68.606 33.827 31.738 1.00 26.13
	ATOM	1708 O ALA 379	68.174 34.580 32.601 1.00 26.46
	ATOM	1709 N CYA 380	69.826 33.935 31.230 1.00 27.61
	ATOM	1710 CA CYA 380	70,742 34,973 31.667 1.00 29.74
5	ATOM	1711 CB CYA 380	72.070 34.865 30.923 1.00 35.44
•	ATOM	1712 SG CYA 380	73.081 33.458 31.417 1.00 42.61
•	ATOM	1713 AS CYA 380	74.829 33.691 29.945 1.00 55.91
	ATOM	1714 C CYA 380	70.142 36.349 31.446 1.00 29.07
	ATOM	1715 O CYA 380	70.243 37.225 32.303 1.00 29.46
10	ATOM	1716 N HIS 381	69.494 36.538 30.304 1.00 28.29
••	ATOM	1717 CA HIS 381	68.885 37.824 30.002 1.00 26.84
	ATOM	1718 CB HIS 381	68.384 37.880 28.557 1.00 23.13
	ATOM	1719 CG HIS 381	67.597 39.113 28.259 1.00 19.84
	ATOM	1720 CD2 HIS 381	67.993 40.365 27.931 1.00 18.68
15	ATOM	1721 ND1 HIS 381	66,229 39.169 28.403 1.00 19.47
13	ATOM	1722 CE1 HIS 381	65.817 40.407 28.190 1.00 18.64
	ATOM	1722 CET HIS 381	66.868 41.149 27.900 1.00 18.29
	ATOM	1724 C HIS 381	67.747 38.157 30.967 1.00 26.78
	ATOM	1725 O HIS 381	67.560 39.314 31.337 1.00 26.39
20	ATOM	1726 N ALA 382	66.964 37.158 31.347 1.00 27.78
20	ATOM	1727 CA ALA 382	65.867 37.395 32.269 1.00 29.45
	ATOM	1728 CB ALA 382	65.077 36.125 32.471 1.00 29.51
	ATOM	1729 C ALA 382	66.425 37.904 33.604 1.00 31.74
	ATOM	1730 O ALA 382	65.932 38.882 34.159 1.00 32.60
25	ATOM	1731 N SER 383	67.483 37.262 34.093 1.00 33.02
	ATOM	1732 CA SER 383	68.109 37.662 35.350 1.00 34.69
	ATOM	1733 CB SER 383	69.212 36.677 35.733 1.00 36.18
	ATOM	1734 OG SER 383	68.663 35.386 35.933 1.00 40.61
	ATOM	1735 C SER 383	68.689 39.064 35.242 1.00 33.49
30	ATOM	1736 O SER 383	68.526 39.889 36.146 1.00 34.28
	ATOM	1737 N ARG 384	69.377 39.332 34.141 1.00 32.60
	ATOM	1738 CA ARG 384	69.955 40.642 33.938 1.00 32.60
	ATOM	1739 CB ARG 384	70.926 40.638 32.762 1.00 33.60
	ATOM	1740 CG ARG 384	71.429 42.013 32.409 1.00 36.33
35	ATOM	1741 CD ARG 384	72.875 41.975 31.993 1.00 39.62
55	ATOM	1742 NE ARG 384	73.760 42.260 33.114 1.00 41.76
	ATOM	1743 CZ ARG 384	74.587 43.301 33.179 1.00 41.92
	ATOM	1744 NH1 ARG 384	
	ATOM	1745 NH2 ARG 384	
40			68.862 41.694 33.758 1.00 32.28
	ATOM	1747 O ARG 384	69.014 42.831 34.213 1.00 33.27
	ATOM	1748 N PHE 385	67.739 41.311 33.159 1.00 29.13
	ATOM	1749 CA PHE 385	66.663 42.259 32.977 1.00 27.55
	ATOM	1750 CB PHE 385	65.552 41.687 32.105 1.00 26.89
45	ATOM	1751 CG PHE 385	64.415 42.641 31.888 1.00 25.11
	ATOM	1752 CD1 PHE 385	64.495 43.630 30.918 1.00 24.94
	ATOM	1752 CD1 PHE 385	63.281 42.580 32.689 1.00 25.01
	ATOM	1754 CE1 PHE 385	63.466 44.547 30.753 1.00 25.50
	ATOM	1755 CE2 PHE 385	62.244 43.495 32.531 1.00 24.06
50	ATOM	1756 CZ PHE 385	62.338 44.482 31.563 1.00 25.44
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    ATOM 1758 O PHE 385
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    ATOM 1761 CB LEU 386
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    ATOM 1762 C LEU 386
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    ATOM 1763 O LEU 386
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    ATOM 1765 CA HIS 387
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    ATOM 1766 CB HIS 387
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    ATOM 1767 CG HIS 387
    ATOM 1768 CD2 HIS 387
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                               71.408 41.161 38.543 1.00 40.97
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    ATOM 1770 CE1 HIS 387
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    ATOM 1771 NE2 HIS 387
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    ATOM 1775 CA MET 388
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    ATOM 1777 CG MET 388
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    ATOM 1778 SD MET 388
ATOM 1779 CE MET 388
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                                68.208 44.370 30.709 1.00 42.36
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                               67.363 48.506 35.886 1.00 49.79
    ATOM 1781 O MET 388
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                               66.163 46.610 36.075 1.00 52.74
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    ATOM 1783 CA LYS 389
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                                                              ALTA
     ATOM 1784 CB LYS 389
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                                                               ALTA
     ATOM 1785 CG LYS 389
                                63.227 46.087 35.161 1.00 57.76
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                                62.029 45.156 35.212 1.00 55.98
                                                               ALTA
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     ATOM 1787 CE LYS 389
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                                                               ALTA
    ATOM 1788 NZ LYS 389
                                61.267 43.040 36.311 1.00 55.55
                                                               ALTA
                               65.177 47.767 38.064 1.00 56.69
     ATOM 1789 C LYS 389
                                                              ALTA
     ATOM 1790 O LYS 389
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                                                              ALTA
                               65.955 47.038 38.839 1.00 55.21
     ATOM 1791 N VAL 390
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     ATOM 1792 CA VAL 390
                                66.225 47.386 40.236 1.00 51.78
                                66.999 46.231 40.985 1.00 50.07
     ATOM 1793 CB VAL 390
                                 67.648 46.726 42.263 1.00 49.74
     ATOM 1794 CG1 VAL 390
                                 66.037 45.093 41.317 1.00 49.06
     ATOM 1795 CG2 VAL 390
                               67.053 48.681 40.227 1.00 49.38
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     ATOM 1796 C VAL 390
                               66.785 49.605 40.992 1.00 48.71
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     ATOM 1798 N GLU 391
     ATOM 1799 CA GLU 391
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     ATOM 1800 CB GLU 391
                                70.156 49.488 38.438 1.00 45.24
                                70.793 48.207 38.997 1.00 47.65
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     ATOM 1801 CG GLU 391
     ATOM 1802 CD GLU 391
                                71.461 48.388 40.358 1.00 50.29
     ATOM 1803 OE1 GLU 391
                                71.141 49.373 41.063 1.00 50.68
     ATOM 1804 OE2 GLU 391
                                72.310 47.535 40.718 1.00 50.85
                               68.324 51.174 38.458 1.00 45.28
     ATOM 1805 C GLU 391
                               68.568 52.286 38.940 1.00 46.46
     ATOM 1806 O GLU 391
50
```

	ATOM	1807 N CYA 392	67.568 51.024 37.372 1.00 43.33
	ATOM	1808 CA CYA 392	67.071 52.192 36.643 1.00 42.28
	ATOM	1809 CB CYA 392	67.519 52.096 35.197 1.00 42.45
	ATOM	1810 SG CYA 392	69.280 52.182 35.127 1.00 43.69
5	ATOM	1811 AS CYA 392	69.908 51.044 33.336 1.00 48.17
	ATOM	1812 C CYA 392	65.589 52.493 36.709 1.00 42.51
	ATOM	1813 O CYA 392	64.792 51.634 37.070 1.00 43.30
	ATOM	1814 N PRO 393	65.205 53.752 36.418 1.00 42.13
	ATOM	1815 CD PRO 393	66.109 54.899 36.199 1.00 40.54
10	ATOM	1816 CA PRO 393	63.794 54.182 36.441 1.00 42.26
	ATOM	1817 CB PRO 393	63.896 55.710 36.365 1.00 41.47
	ATOM	1818 CG PRO 393	65.189 55.938 35.614 1.00 41.10
	ATOM	1819 C PRO 393	62.954 53.606 35.281 1.00 43.20
	ATOM	1820 O PRO 393	63.463 53.452 34.163 1.00 42.61
15	ATOM	1821 N THR 394	61.686 53.305 35.559 1.00 43.70
13	ATOM	1822 CA THR 394	60.764 52.755 34.564 1.00 45.50
	ATOM	1823 CB THR 394	59.340 52.609 35.129 1.00 47.20
	ATOM	1824 OG1 THR 394	59.304 53.139 36.464 1.00 50.57
	ATOM	1825 CG2 THR 394	58.878 51.150 35.137 1.00 47.99
20	ATOM	1826 C THR 394	60.682 53.583 33.283 1.00 44.58
20	ATOM	1827 O THR 394	60.409 53.054 32.215 1.00 46.36
	ATOM	1828 N GLU 395	60.899 54.888 33.396 1.00 42.88
	ATOM	1829 CA GLU 395	60.842 55.790 32.246 1.00 40.54
	ATOM	1830 CB GLU 395	61.096 57.234 32.699 1.00 40.69
25	ATOM	1831 C GLU 395	61.799 55.421 31.098 1.00 38.51
23	ATOM	1832 O GLU 395	61.628 55.877 29.968 1.00 39.41
	ATOM	1833 N LEU 396	62.828 54.640 31.402 1.00 35.60
	ATOM	1834 CA LEU 396	63.795 54.220 30.386 1.00 33.11
	ATOM	1835 CB LEU 396	65.169 54.003 31.027 1.00 33.60
30	ATOM	1836 CG LEU 396	65.831 55.230 31.660 1.00 34.54
	ATOM	1837 CD1 LEU 396	67.160 54.835 32.282 1.00 32.83
	ATOM	1838 CD2 LEU 396	66.026 56.308 30.599 1.00 35.71
	ATOM	1839 C LEU 396	63.388 52.940 29.660 1.00 30.95
	ATOM	1840 O LEU 396	63,950 52.605 28.624 1.00 30.90
35	ATOM	1841 N PHE 397	62,422 52,227 30,223 1,00 30,18
	ATOM	1842 CA PHE 397	61.961 50.970 29.654 1.00 28.80
	ATOM	1843 CB PHE 397	61.712 49.946 30.777 1.00 28.10
	ATOM	1844 CG PHE 397	62.938 49.604 31.592 1.00 28.96
	ATOM	1845 CD1 PHE 397	63.403 50.472 32.591 1.00 28.39
40	ATOM	1846 CD2 PHE 397	63.636 48.422 31.359 1.00 26.28
	ATOM	1847 CE1 PHE 397	64.546 50.166 33.337 1.00 28.44
	ATOM	1848 CE2 PHE 397	64.784 48.107 32.103 1.00 29.21
	ATOM	1849 CZ PHE 397	65,240 48,984 33,096 1,00 27,37
	ATOM	1850 C PHE 397	60.683 51.093 28.836 1.00 27.54
45	ATOM	1851 O PHE 397	59.630 51.431 29.370 1.00 26.96
-	ATOM	1852 N PRO 398	60.753 50.836 27.501 1.00 27.41
	ATOM	1853 CD PRO 398	61.968 50.600 26.686 1.00 25.42
	ATOM	1854 CA PRO 398	59.560 50.920 26.654 1.00 25.90
	ATOM	1855 CB PRO 398	60.068 50.383 25.320 1.00 25.26
50	ATOM	1856 CG PRO 398	61.490 50.893 25.290 1.00 23.99

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ATOM 1857 C PRO 398
                               58.494 49.995 27.272 1.00 25.86
                               58.839 48.962 27.843 1.00 25.82
    ATOM 1858 O PRO 398
                               57.197 50.355 27.175 1.00 25.52
    ATOM 1859 N PRO 399
                                56.627 51.576 26.578 1.00 25.49
    ATOM 1860 CD PRO 399
    ATOM 1861 CA PRO 399
                                56.145 49.510 27.754 1.00 25.42
                                54.861 50.181 27.273 1.00 26.23
    ATOM 1862 CB PRO 399
                                55.237 51.609 27.156 1.00 25.25
    ATOM 1863 CG PRO 399
    ATOM 1864 C PRO 399
                               56.198 48.043 27.317 1.00 26.08
                               56.132 47.131 28.159 1.00 25.45
    ATOM 1865 O PRO 399
                               56.350 47.810 26.019 1.00 25.57
10
    ATOM 1866 N LEU 400
    ATOM 1867 CA LEU 400
                                56.406 46.440 25.509 1.00 26.27
                                56.404 46.418 23.980 1.00 25.03
    ATOM 1868 CB LEU 400
    ATOM 1869 CG LEU 400
                                56.117 45.042 23.363 1.00 24.51
    ATOM 1870 CD1 LEU 400
                                54.757 44.530 23.806 1.00 23.22
     ATOM 1871 CD2 LEU 400
                                56.173 45.149 21.862 1.00 23.70
15
                               57.602 45.657 26.067 1.00 27.06
     ATOM 1872 C LEU 400
                               57.484 44.465 26.363 1.00 27.41
    ATOM 1873 O LEU 400
                               58.736 46.339 26.231 1.00 27.16
     ATOM 1874 N PHE 401
                                59.966 45.754 26.779 1.00 27.06
     ATOM 1875 CA PHE 401
                                61.047 46.833 26.802 1.00 26.60
20
     ATOM 1876 CB PHE 401
     ATOM 1877 CG PHE 401
                                62.408 46.351 27.217 1.00 28.08
                                62.918 45.138 26,747 1.00 27.45
     ATOM 1878 CD1 PHE 401
     ATOM 1879 CD2 PHE 401
                                63.223 47.165 28.013 1.00 27.48
                                64.220 44.746 27.055 1.00 26.95
     ATOM 1880 CE1 PHE 401
     ATOM 1881 CE2 PHE 401
                                64.523 46.786 28.327 1.00 27.97
25
     ATOM 1882 CZ PHE 401
                                65.028 45.575 27.846 1.00 28.46
                               59.690 45.247 28.205 1.00 27.62
     ATOM 1883 C PHE 401
     ATOM 1884 O PHE 401
                               60.046 44.125 28.570 1.00 26.24
                               59.036 46.082 29.002 1.00 28.75
     ATOM 1885 N LEU 402
     ATOM 1886 CA LEU 402
                                58.692 45.719 30.366 1.00 29.58
30
     ATOM 1887 CB LEU 402
                                58.064 46.910 31.088 1.00 30.04
     ATOM 1888 CG LEU 402
                                59.025 47.974 31.594 1.00 30.14
     ATOM 1889 CD1 LEU 402
                                58.270 49.263 31.880 1.00 29.61
                                59.734 47.438 32.827 1.00 27.99
     ATOM 1890 CD2 LEU 402
     ATOM 1891 C LEU 402
                               57.693 44.583 30.368 1.00 30.10
35
                               57.836 43.631 31.121 1.00 29.78
     ATOM 1892 O LEU 402
                               56.688 44.683 29.510 1.00 30.49
     ATOM 1893 N GLU 403
                                55.646 43.671 29.453 1.00 32.60
     ATOM 1894 CA GLU 403
                                54.562 44.094 28.469 1.00 37.01
     ATOM 1895 CB GLU 403
     ATOM 1896 CG GLU 403
                                53.329 43.218 28.520 1.00 44.01
40
                                52.263 43.632 27.523 1.00 48.50
     ATOM 1897 CD GLU 403
                                52.516 44.525 26.677 1.00 49.66
     ATOM 1898 OE1 GLU 403
     ATOM 1899 OE2 GLU 403
                                 51.157 43.050 27.594 1.00 53.06
                               56.083 42.237 29.151 1.00 32.03
     ATOM 1900 C GLU 403
                                55.627 41.304 29.816 1.00 32.58
     ATOM 1901 O GLU 403
45
                                56.955 42.078 28.159 0.50 31.51
                                                              ALTA
     ATOM 1902 N VAL 404
                                57.450 40.765 27.739 0.50 30.96
                                                               ALTA
     ATOM 1903 CA VAL 404
     ATOM 1904 CB VAL 404
                                58.108 40.849 26.333 0.50 30.32
                                                               ALTA
                                 58.616 39.489 25.889 0.50 28.72
     ATOM 1905 CG1 VAL 404
                                                               ALTA
                                57.115 41.388 25.328 0.50 31.67
     ATOM 1906 CG2 VAL 404
                                                               ALTA
50
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ALTA
    ATOM 1907 C VAL 404
                               58.465 40.149 28.696 0.50 30.45
                               58.549 38.926 28.822 0.50 30.10
                                                              ALTA
     ATOM 1908 O VAL 404
                               59,224 41.002 29.369 1.00 30.16
    ATOM 1909 N PHE 405
                                60.266 40.549 30.263 1.00 30.65
    ATOM 1910 CA PHE 405
    ATOM 1911 CB PHE 405
                                61.577 41.221 29.863 1.00 28.92
                                62.062 40.834 28.493 1.00 26.31
    ATOM 1912 CG PHE 405
                                62.342 41.804 27.543 1.00 25.72
    ATOM 1913 CD1 PHE 405
                                62.269 39.500 28.166 1.00 25.92
    ATOM 1914 CD2 PHE 405
                                62.827 41.456 26.278 1.00 26.78
    ATOM 1915 CE1 PHE 405
                                62.752 39.139 26.910 1.00 25.39
10
    ATOM 1916 CE2 PHE 405
                                63.034 40.122 25.962 1.00 24.39
    ATOM 1917 CZ PHE 405
    ATOM 1918 C PHE 405
                               60.011 40.674 31.771 1.00 32.10
     ATOM 1919 O PHE 405
                               60.903 40.237 32.533 1.00 33.88
                                 58.936 41.169 32.188 1.00 34.95
    ATOM 1920 OXT PHE 405
                               67.542 37.066 11.311 1.00 26.83
    ATOM
             1 O1 HOH 501
15
                               68.713 41.227 12.821 1.00 23.42
    ATOM
             3 O1 HOH 502
    ATOM
             2 O1 HOH 503
                               64.446 40.325 12.123 1.00 22.84
             4 O1 HOH 504
                               62.236 39.752 15.941 1.00 17.97
    ATOM
             5 O1 HOH 505
                               48.732 20.137 5.515 1.00 50.48
    ATOM
             6 O1 HOH 506
                               47.365 21.522 3.716 1.00 53.40
20
    ATOM
             7 O1 HOH 507
                               50.211 23.203 7.900 1.00 32.66
     ATOM
                               51.043 20.258 8.253 1.00 21.81
     ATOM
             8 O1 HOH 508
             9 O1 HOH 509
                               48.225 18.176 7.905 1.00 38.96
     ATOM
                              49.569 20.871 11.586 1.00 32.97
     ATOM
             10 O1 HOH 510
                               53.732 17.159 10.856 1.00 47.20
             11 O1 HOH 511
25
     ATOM
     ATOM
             12 O1 HOH 512
                               56.201 16.223 12.164 1.00 18.50
                               56.653 12.298 10.528 1.00 27.71
     ATOM
             13 O1 HOH 513
             14 O1 HOH 514
                               58.661 10.694 9.014 1.00 46.73
     ATOM
                               62.950 10.692 11.952 1.00 43.05
     ATOM
             15 O1 HOH 515
                               66.411 11.552 10.897 1.00 37.36
30
    ATOM
             16 O1 HOH 516
                               68.949 13.188 12.029 1.00 39.28
    ATOM
             17 O1 HOH 517
                               71.997 15.171 8.362 1.00 49.69
             18 O1 HOH 518
     ATOM
             19 O1 HOH 519
                               71.946 17.928 6.743 1.00 24.50
     ATOM
             20 O1 HOH 520
                               75.117 15.684 9.377 1.00 35.98
     ATOM
                               76.677 12.815 10.294 1.00 49.33
             21 O1 HOH 521
35
     ATOM
                               81.421 15.415 15.139 1.00 46.74
     ATOM
             22 O1 HOH 522
                               78.784 21.696 17.564 1.00 49.01
             23 O1 HOH 523
     ATOM
                               79.954 24.822 17.152 1.00 42.91
     ATOM
             24 O1 HOH 524
                               82.199 30.253 18.821 1.00 40.27
     ATOM
             25 O1 HOH 525
                               82.862 33.444 21.988 1.00 46.81
40
    ATOM
             26 O1 HOH 526
             27 O1 HOH 527
                               76.608 30.793 23.452 1.00 46.22
     ATOM
                               74.726 30.483 25.469 1.00 43.76
     ATOM
             28 O1 HOH 528
     ATOM
             29 O1 HOH 529
                               77.059 28.762 20.900 1.00 33.67
             30 O1 HOH 530
                               75.935 33.279 12.269 1.00 25.26
     ATOM
                               77.402 34.447 10.087 1.00 37.04
     ATOM
             31 O1 HOH 531
                               74.054 29.941 9.998 1.00 26.86
     ATOM
             32 O1 HOH 532
                               69.544 32.658 7.572 1.00 40.34
     ATOM
             33 O1 HOH 533
                               66.709 33.618 8.477 1.00 20.63
     ATOM
             34 O1 HOH 534
                               68.073 35.828 8.931 1.00 23.99
     ATOM
             35 O1 HOH 535
                               61.865 45.643 14.011 1.00 40.43
50
     ATOM
             36 O1 HOH 536
```

	ATOM	37 O1 HO	H 537	63.662	46.881	15.670	1.00 28.04
	ATOM	38 O1 HO	H 538	63.391	49.310	13.883	1.00 39.59
	ATOM	39 O1 HO	H 539	63.491	50.570	10.631	1.00 52.34
	ATOM	40 O1 HO	H 540	64.592	46.849	10.299	1.00 26.63
5	ATOM	41 O1 HO	H 541	55.575	41.632	10.980	1.00 38.06
	ATOM	42 O1 HO	H 542	51.631	42.062	17.343	1.00 45.99
	ATOM	43 O1 HO	H 543	52.755	43.156	20.209	1.00 34.17
	ATOM	44 O1 HO		57.061	49.627	24.004	1.00 24.09
	ATOM	45 O1 HO	H 545	61.040	50.561	21.351	1.00 30.91
10	ATOM	46 O1 HO			53.616	18.390	1.00 30.91
	ATOM	47 O1 HO			58.813	29.014	1.00 59.25
	ATOM	48 O1 HO			52.905	31.175	1.00 40.12
	ATOM	49 O1 HO			50.496	37.543	1.00 52.28
	ATOM	50 O1 HO			49.704	40.891	1.00 54.18
15	ATOM	51 O1 HO			46.430	40.384	1.00 43.84
13	ATOM	52 O1 HO			14.459	39.268	1.00 44.73
	ATOM	53 O1 HO			43.920	33.936	1.00 42.88
	ATOM	54 O1 HO			39.071	34.046	1.00 45.07
	ATOM	55 O1 HO			36.925	28.676	1.00 45.07
20	ATOM	56 O1 HO				26.610	1.00 23.30
20	ATOM	57 O1 HO			33.760	25.609	1.00 21.14
	ATOM	58 O1 HO			29.906	24.568	1.00 55.05
	ATOM	59 O1 HO			27.540	24.855	1.00 37.37
	ATOM	60 O1 HO			28.785	27.536	1.00 59.05
25	ATOM	61 O1 HO			30.642	28.821	1.00 54.44
23	ATOM	62 O1 HO			24.645	32.964	1.00 50.75
	ATOM	63 O1 HO			21.149	28.711	1.00 50.75
	ATOM	64 O1 HO				26.900	1.00 51.41
	ATOM	65 O1 HO			20.422	23.303	1.00 32.30
30					18.167	23.386	1.00 40.32
30	ATOM				22.649	25.573	1.00 63.08
	ATOM ATOM					25.147	1.00 48.83
		68 O1 HO			21.446	28.543	1.00 43.39
	ATOM	69 O1 HO			20.179		
25	ATOM	70 O1 HO			23.216	30.984 32.696	1.00 50.28 1.00 43.96
35	ATOM	71 O1 HO			24.880		
	ATOM	72 O1 HO			29.683	30.486	1.00 44.51
	ATOM	73 O1 HO				30.521	1.00 36.28
	ATOM	74 O1 HO				29.178	1.00 51.45
4.0	ATOM	75 O1 HO				30.420	1.00 31.03
40	ATOM	76 O1 HO					1.00 19.54
	ATOM	77 O1 HO				28.703	1.00 33.04
	ATOM	78 O1 HO			15.471		1.00 35.68
	ATOM	79 O1 HO					1.00 50.46
	ATOM	80 O1 HO		53.726			1.00 41.44
45	ATOM	81 O1 HO		57.223			1.00 48.31
	ATOM	82 O1 HO		61.169			1.00 17.60
	ATOM	83 O1 HO					1.00 23.93
	ATOM	84 O1 HO		67.033			1.00 26.21
	ATOM	85 O1 HO					1.00 59.67
50	ATOM	86 O1 HO	H 586	68.489 2	22.464	0.350	1.00 37.85

WO 99/26966

```
65,794 23.354 0.823 1.00 27.38
             87 O1 HOH 587
    ATOM
                               67.550 26.810 0.937 1.00 37.18
             88 O1 HOH 588
    ATOM
                                64.646 28.208 3.323 1.00 36.74
             89 O1 HOH 589
    ATOM
                               67.215 31.103 3.174 1.00 30.29
             90 O1 HOH 590
     ATOM
                                64.164 35.667 6.220 1.00 39.72
             91 O1 HOH 591
5
    ATOM
                                62.810 37.518 4.836 1.00 48.48
             92 O1 HOH 592
     ATOM
                                68.105 36.898 6.110 1.00 58.00
             93 O1 HOH 593
     ATOM
                                57,390 37,485 2,631 1,00 37,29
     ATOM
             94 O1 HOH 594
                                53.088 36.068 3.949 1.00 50.10
    ATOM
             95 O1 HOH 595
                                52.974 34.676 6.758 1.00 42.52
    ATOM
             96 O1 HOH 596
10
                                58.581 31.465 2.076 1.00 32.18
     ATOM
             97 O1 HOH 597
                                52.786 23.277 1.357 1.00 28.98
     ATOM
             98 O1 HOH 598
                                47.501 26.551 7.672 1.00 47.83
             99 O1 HOH 599
     ATOM
                                46.411 35.754 14.049 1.00 53.46
             100 O1 HOH 600
     ATOM
                                63.514 14.944 15.842 1.00 55.02
            101 O1 HOH 601
     ATOM
15
                                67.943 11.792 3.438 1.00 61.21
            102 O1 HOH 602
     ATOM
                                62.232 9.378 3.311 1.00 35.65
            103 O1 HOH 603
     ATOM
                                76.734 22.468 5.002 1.00 42.56
            104 O1 HOH 604
     ATOM
                                83.589 28.967 9.626 1.00 50.64
            105 O1 HOH 605
     ATOM
                                82.807 43.437 17.940 1.00 39.28
            106 O1 HOH 606
     ATOM
20
            107 O1 HOH 607
                                83.882 45.673 20.638 1.00 41.64
     ATOM
                                80.215 41.021 23.441 1.00 43.16
            108 O1 HOH 608
     ATOM
                                79.459 46.296 31.165 1.00 32.40
            109 O1 HOH 609
     ATOM
                                81.880 47.681 33.923 1.00 46.96
            110 O1 HOH 610
     ATOM
                                75.594 46.142 30.384 1.00 28.64
25
     ATOM
            111 O1 HOH 611
            112 O1 HOH 612
                                77.118 40.568 32.575 1.00 34.21
     ATOM
                                73.563 41.750 36.926 1.00 26.07
            113 O1 HOH 613
     ATOM
                                75.955 56.565 28.863 1.00 46.31
             114 O1 HOH 614
     ATOM
                                79.915 59.136 15.809 1.00 50.81
             115 O1 HOH 615
     ATOM
                                77.390 52.542 8.816 1.00 34.34
            116 O1 HOH 616
30
     ATOM
            117 O1 HOH 617
                                72.726 25.005 29.671 1.00 62.84
     ATOM
                                52.664 40.106 24.800 1.00 46.39
     ATOM 2038 C ACY 701
                                53.721 39.649 24.298 1.00 47.12
     ATOM 2039 O ACY 701
                                  51.652 40.521 24.172 1.00 46.96
     ATOM 2040 OXT ACY 701
     ATOM 2041 CH3 ACY 701
                                  52.600 40.162 26.329 1.00 45.99
35
                              66.961 42.243 18.491 1.00 22.34
     ATOM 2050 C1 T3
                          1
                              68.748 43.593 23.015 1.00 21.84
     ATOM 2051 C2 T3
                              66.873 43.557 18.970 1.00 23.43
     ATOM 2052 C3 T3
                              69.252 44.540 23.871 1.00 22.31
     ATOM 2053 C4 T3
     ATOM 2054 C5 T3
                              67.638 43.989 20.011 1.00 24.83
40
                          1
                              68.851 44.553 25.178 1.00 25.16
     ATOM 2055 C6 T3
                          1
     ATOM 2056 C7 T3
                              68.541 43.108 20.632 1.00 24.65
                          1
                              67.895 43.567 25.639 1.00 21.93
     ATOM 2057 C8 T3
                          1
                              68.665 41.792 20.183 1.00 25.09
     ATOM 2058 C9 T3
                          1
                               67,427 42.654 24.733 1.00 23.66
     ATOM 2059 C10 T3
45
                           1
                               67.878 41.380 19.117 1.00 23.12
     ATOM 2060 C11 T3
                           1
                               67.829 42.624 23.384 1.00 19.67
     ATOM 2061 C12 T3
                           1
                               66.055 41.788 17.371 1.00 18.97
     ATOM 2062 C13 T3
                           1
                               66.721 40.956 16.295 1.00 19.32
     ATOM 2063 C15 T3
                           1
                               65.901 40.829 15.051 1.00 19.02
50
     ATOM 2064 C17 T3
                           1
```

	ATOM	2065 II T3	1	67.393 45.986 20.621 1.00 25.29
	ATOM	2066 I2 T3	1	69.483 46.066 26.432 1.00 26.49
	ATOM	2067 I3 T3	1	70.019 40.450 20.975 1.00 25.67
	ATOM	2068 N1 T3	1	68.131 41.337 16.037 1.00 15.12
5	ATOM	2069 O1 T3	1	67.542 43.587 26.966 1.00 21.79
	ATOM	2070 O2 T3	1	69.259 43.600 21.682 1.00 22.05
	ATOM	2071 O3 T3	1	66.504 40.852 13.963 1.00 20.38
	<b>ATOM</b>	2072 O4 T3	1	64.675 40.731 15.192 1.00 20.16
	END			

#### **APPENDIX 7**

#### TRBTRIAC.PDB

REMARK TR-beta Triac Full length numbering

REMARK refinement resolution: 100 - 2.9 A r= 0.273258 free\_r= 0.333794

5 REMARK wa= 5.78307

REMARK target= mlf cycles= 1 steps= 25

REMARK a= 68.72 b= 68.72 c= 130.092 alpha= 90 beta= 90 gamma= 120

REMARK ncs= none

REMARK initial B-factor correction: "none"

10 REMARK ALA 199 to ALA 201 from His-tag

REMARK

REMARK Four cacodylate-modified cysteines (CYA)

REMARK Cys294, Cys298, Cys388, Cys434

REMARK cacodylate modeled as single arsenic atom

15 REMARK

REMARK side chain of certain residues modeled as ALA due to poor density;

REMARK however, residue name reflects true residue for clarity

**REMARK** 

REMARK amino acid sequence confirmed,

20 REMARK differing from that reported by Weinberger et. al.

REMARK in the following codons:

REMARK 243 Pro - Arg

REMARK 337 lle - Thr

REMARK 451 Leu - Phe

25 REMARK as reported by Sakurai et. al.

REMARK note also correction of initiation codon,

REMARK yielding a polypeptide of 461 amino acids

JRNL AUTH A.SAKURAI, A.NAKAI, L.J. DEGROOT

JRNL TITL STRUCTURAL ANALYSIS OF HUMAN THYROID HORMONE

30 RECEPTOR

JRNL TITL2 BETA GENE

JRNL REF MOL.CELL.ENDO. V.71 1990

JRNL AUTH

C.WEINBERGER,

C.C.THOMPSON,R.LEBO,D.J.GRUOL,R.M.EVANS

35 JRNL TITL THE C-ERB-A GENE ENCODES A THYROID HORMONE RECEPTOR

JRNL REF NATURE

V.324 6098 1986

ATOM	1 CB ALA 199	31.247 28.289 43.613 1.00 71.30	PROT
ATOM	2 C ALA 199	32.916 26.485 44.170 1.00 68.99	PROT
ATOM	3 O ALA 199	33.485 25.410 43.976 1.00 63.84	PROT
ATOM	4 N ALA 199	30.462 25.993 44.096 1.00 75.00	PROT
<b>ATOM</b>	5 CA ALA 199	31.571 26.795 43.497 1.00 73.24	PROT
ATOM	6 N ALA 200	33.419 27.432 44.958 1.00 73.81	PROT
ATOM	7 CA ALA 200	34.686 27.251 45.658 1.00 67.87	PROT
ATOM	8 CB ALA 200	35.182 28.583 46.203 1.00 62.83	PROT
ATOM	9 C ALA 200	34.539 26.239 46.791 1.00 63.23	PROT
ATOM	10 O ALA 200	35.486 25.986 47.534 1.00 59.14	PROT
	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	ATOM 2 C ALA 199 ATOM 3 O ALA 199 ATOM 4 N ALA 199 ATOM 5 CA ALA 199 ATOM 6 N ALA 200 ATOM 7 CA ALA 200 ATOM 8 CB ALA 200 ATOM 9 C ALA 200	ATOM 2 C ALA 199 32.916 26.485 44.170 1.00 68.99 ATOM 3 O ALA 199 33.485 25.410 43.976 1.00 63.84 ATOM 4 N ALA 199 30.462 25.993 44.096 1.00 75.00 ATOM 5 CA ALA 199 31.571 26.795 43.497 1.00 73.24 ATOM 6 N ALA 200 33.419 27.432 44.958 1.00 73.81 ATOM 7 CA ALA 200 34.686 27.251 45.658 1.00 67.87 ATOM 8 CB ALA 200 35.182 28.583 46.203 1.00 62.83 ATOM 9 C ALA 200 34.539 26.239 46.791 1.00 63.23

	4 TO 3 4	11 N. A.T.A. 201	33.345 25.670 46.932 1.00 56.98	PROT
	ATOM	11 N ALA 201 12 CA ALA 201	33.343 23.670 46.932 1.00 30.98	PROT
	ATOM ATOM	13 CB ALA 201	31.776 23.992 47.744 1.00 40.35	PROT
	ATOM		34.248 23.662 47.762 1.00 53.15	PROT
5	ATOM	15 O ALA 201	34.624 22.938 48.679 1.00 54.90	PROT
3	ATOM	16 N GLU 202	34.789 23.645 46.546 1.00 44.13	PROT
	ATOM	17 CA GLU 202	35.891 22.767 46.190 1.00 37.47	PROT
	ATOM	18 CB GLU 202	36.086 22.760 44.671 1.00 37.74	PROT
	ATOM	19 CG GLU 202	37.060 21.702 44.173 1.00 57.14	PROT
10	ATOM	20 CD GLU 202	36.457 20.303 44.140 1.00 61.74	PROT
10	ATOM	21 OE1 GLU 202	35.211 20.175 44.133 1.00 63.81	PROT
	ATOM	22 OE2 GLU 202	37.236 19.327 44.115 1.00 65.54	PROT
	ATOM		37.156 23.266 46.878 1.00 35.54	PROT
	ATOM	24 O GLU 202	37.874 22.492 47.510 1.00 32.70	PROT
15	ATOM	25 N GLU 203	37.415 24.566 46.755 1.00 31.79	PROT
13	ATOM	26 CA GLU 203	38.588 25.188 47.366 1.00 33.63	PROT
	ATOM	27 CB GLU 203	38.603 26.683 47.079 1.00 28.28	PROT
	ATOM		38.588 24.948 48.869 1.00 33.86	PROT
	ATOM	29 O GLU 203	39.644 24.818 49.485 1.00 33.10	PROT
20	ATOM		37.393 24.898 49.451 1.00 34.15	PROT
	ATOM	31 CA LEU 204	37.244 24.650 50.876 1.00 33.22	PROT
	ATOM	32 CB LEU 204	35.853 25.081 51.353 1.00 30.47	PROT
	ATOM	33 CG LEU 204	35,567 25.083 52.862 1.00 23.17	PROT
	ATOM	34 CD1 LEU 204	35.904 26.439 53.443 1.00 5.41	PROT
25	ATOM	35 CD2 LEU 204	34.106 24.748 53.111 1.00 12.70	PROT
	ATOM	36 C LEU 204	37.424 23.156 51.100 1.00 40.17	PROT
	ATOM	37 O LEU 204	38.219 22.736 51.951 1.00 45.33	PROT
	ATOM	38 N GLN 205	36.682 22.360 50.329 1.00 43.86	PROT
	ATOM	39 CA GLN 205	36.754 20.899 50.415 1.00 43.96	PROT
30	ATOM	40 CB GLN 205	36.089 20.261 49.184 1.00 45.56	PROT
	ATOM	41 CG GLN 205	34.562 20.195 49.245 1.00 42.39	PROT
	ATOM	42 CD GLN 205	34.022 18.775 49.159 1.00 46.79	PROT
	ATOM	43 OE1 GLN 205	33.258 18.444 48.252 1.00 38.84	PROT
	ATOM	44 NE2 GLN 205	34.412 17.932 50.109 1.00 37.95	PROT
35	ATOM		38.224 20.482 50.483 1.00 42.39	PROT
	ATOM	46 O GLN 205	38.630 19.702 51.355 1.00 36.27	PROT
	ATOM		39.014 21.015 49.553 1.00 42.37	PROT
	ATOM	48 CA LYS 206	40.440 20.729 49.505 1.00 44.40	PROT
40	ATOM	49 CB LYS 206	41.110 21.531 48.385 1.00 38.73	PROT
40	ATOM		41.024 21.118 50.853 1.00 42.36	PROT
	ATOM		41.550 20.271 51.570 1.00 46.93	PROT
	ATOM		40.913 22.401 51.192 1.00 34.68	PROT
	ATOM	53 CA SER 207	41.415 22.933 52.455 1.00 29.43	PROT
15	ATOM	54 CB SER 207 55 OG SER 207	40.690 24.228 52.791 1.00 24.63 41.327 25.332 52.173 1.00 36.56	PROT PROT
45	ATOM		41.254 21.958 53.614 1.00 29.20	
	ATOM		41.254 21.958 53.614 1.00 29.20 42.223 21.623 54.293 1.00 31.01	PROT PROT
	ATOM ATOM			PROT
	ATOM		39.777 20.568 54.928 1.00 27.93	PROT
50	ATOM		38.267 20.216 55.027 1.00 27.93	PROT
50	AIOM	00 CD ILL 200	30.207 20.210 33.027 1.00 37.03	. 1.01

	ATOM	61 CG2 ILE 208	38.062 18.895 55.769 1.00 32.13	PROT
	ATOM	62 CG1 ILE 208	37.528 21.340 55.753 1.00 37.63	PROT
	ATOM	63 CD1 ILE 208	36.788 22.296 54.827 1.00 41.47	PROT
_	ATOM	64 C ILE 208		PROT
5	ATOM	65 O ILE 208	40.905 18.580 55.679 1.00 40.00	PROT
	ATOM	66 N GLY 209	40.928 19.002 53.475 1.00 35.05	PROT
	ATOM	67 CA GLY 209	41.698 17.809 53.181 1.00 31.94	PROT
	ATOM	68 C GLY 209	40.826 16.695 52.643 1.00 28.66	PROT
	ATOM	69 O GLY 209	41.257 15.553 52.532 1.00 19.46	PROT
10	ATOM	70 N HIS 210	39.586 17.021 52.313 1.00 20.47	PROT
	ATOM	71 CA HIS 210	38.684 16.018 51.774 1.00 26.99	PROT
	ATOM	72 CB HIS 210	37.240 16.451 52.012 1.00 37.16	PROT
	ATOM	73 C HIS 210	38.959 15.806 50.266 1.00 27.75	PROT
	ATOM	74 O HIS 210	39.328 16.741 49.550 1.00 34.08	PROT
15	ATOM	75 N LYS 211	38.807 14.566 49.805 1.00 16.50	PROT
	ATOM	76 CA LYS 211	39.019 14.206 48.403 1.00 5.57	PROT
	ATOM	77 CB LYS 211	39.932 12.981 48.295 1.00 5.67	PROT
	ATOM	78 CG LYS 211	41.370 13.208 48.742 1.00 7.30	PROT
	ATOM	79 CD LYS 211	41.873 14.594 48.347 1.00 14.34	PROT
20	ATOM	80 CE LYS 211	43.339 14.556 47.897 1.00 29.48	PROT
	ATOM	81 NZ LYS 211	43.777 15.851 47.262 1.00 33.43	PROT
	ATOM	82 C LYS 211	37.642 13.861 47.876 1.00 2.73	PROT
	ATOM	83 O LYS 211	37.176 12.741 48.039 1.00 6.57	PROT
	ATOM	84 N PRO 212	36.983 14.813 47.208 1.00 2.00	PROT
25	ATOM	85 CD PRO 212	37.472 16.156 46.846 1.00 10.43	PROT
	ATOM	86 CA PRO 212	35.642 14.542 46.689 1.00 2.05	PROT
	ATOM	87 CB PRO 212	35.088 15.928 46.341 1.00 10.09	PROT
	ATOM	88 CG PRO 212	36.240 16.888 46.422 1.00 8.43	PROT
	ATOM	89 C PRO 212	35.523 13.578 45.520 1.00 2.00	PROT
30	ATOM	90 O PRO 212	36.344 13.554 44.611 1.00 6.04	PROT
	ATOM	91 N GLU 213	34.476 12.773 45.577 1.00 2.68	PROT
	ATOM	92 CA GLU 213	34.181 11.817 44.542 1.00 6.81	PROT
	ATOM	93 CB GLU 213	33.539 10.594 45.173 1.00 7.20	PROT
	ATOM	94 CG GLU 213	34.222 10.232 46.462 1.00 15.33	PROT
35	ATOM	95 CD GLU 213	34.293 8.743 46.689 1.00 21.36	PROT
	ATOM	96 OE1 GLU 213	33.334 8.051 46.290 1.00 29.32	PROT
	ATOM	97 OE2 GLU 213	35.301 8.265 47.268 1.00 28.50	PROT
	ATOM	98 C GLU 213	33.229 12.543 43.584 1.00 12.00	PROT
	ATOM	99 O GLU 213	32.693 13.599 43.926 1.00 19.02	PROT
40	ATOM	100 N PRO 214	33.011 11.985 42.375 1.00 25.74	PROT
	ATOM	101 CD PRO 214	33.592 10.692 41.973 1.00 28.98	PROT
	ATOM	102 CA PRO 214	32.145 12.536 41.322 1.00 23.38	PROT
	ATOM	103 CB PRO 214	32.180 11.476 40.232 1.00 18.01	PROT
	ATOM	104 CG PRO 214	33.376 10.665 40.514 1.00 27.50	PROT
45	ATOM	105 C PRO 214	30.715 12.828 41.734 1.00 25.02	PROT
	ATOM	106 O PRO 214	30.069 11.986 42.355 1.00 31.17	PROT
	ATOM	107 N THR 215	30.211 14.009 41.377 1.00 19.56	PROT
	ATOM	108 CA THR 215	28.830 14.352 41.714 1.00 24.48	PROT
	ATOM	109 CB THR 215	28.535 15.841 41.522 1.00 27.13	PROT
50	ATOM	110 OG1 THR 215	5 27.939 16.038 40.234 1.00 40.19	PROT

	ATOM	111 CG2 THR 215	29.805 16.659 41.640 1.00 30.81	PROT
	ATOM	112 C THR 215	27.899 13.562 40.805 1.00 22.14	PROT
	ATOM	113 O THR 215	28.357 12.905 39.883 1.00 27.52	PROT
	ATOM	114 N ASP 216	26.599 13.617 41.072 1.00 35.65	PROT
5	ATOM	115 CA ASP 216	25.631 12.890 40.258 1.00 41.16	PROT
,	ATOM	116 CB ASP 216	24.219 13.091 40.810 1.00 38.17	PROT
	ATOM	117 C ASP 216	25.714 13.370 38.810 1.00 40.44	PROT
	ATOM	118 O ASP 216	25.683 12.569 37.874 1.00 38.26	PROT
	ATOM	119 N GLU 217	25.832 14.682 38.635 1.00 40.14	PROT
10	ATOM	120 CA GLU 217	25.932 15.275 37.305 1.00 38.89	PROT
10	ATOM	121 CB GLU 217	25.883 16.796 37.413 1.00 29.95	PROT
	ATOM	122 C GLU 217	27.231 14.829 36.619 1.00 39.44	PROT
	ATOM	123 O GLU 217	27.245 14.525 35.425 1.00 40.08	PROT
	ATOM	124 N GLU 218	28.319 14.794 37.384 1.00 34.92	PROT
15	ATOM	125 CA GLU 218	29.615 14.370 36.871 1.00 23.70	PROT
13	ATOM	126 CB GLU 218	30.698 14.606 37.924 1.00 18.47	PROT
	ATOM	127 CG GLU 218	30.990 16.067 38.198 1.00 15.66	PROT
	ATOM	128 CD GLU 218	32.085 16.264 39.231 1.00 26.88	PROT
	ATOM	129 OE1 GLU 218	32.164 15.458 40.191 1.00 25.07	PROT
20	ATOM	130 OE2 GLU 218	32.864 17.232 39.078 1.00 33.79	PROT
	ATOM	131 C GLU 218	29.589 12.892 36.491 1.00 21.05	PROT
	ATOM	132 O GLU 218	30.182 12.490 35.495 1.00 24.30	PROT
	ATOM	133 N TRP 219	28.907 12.080 37.288 1.00 13.98	PROT
	ATOM	134 CA TRP 219	28.829 10.660 37.000 1.00 17.30	PROT
25	ATOM	135 CB TRP 219	28.052 9.921 38.089 1.00 16.27	PROT
	ATOM	136 CG TRP 219	28.890 9.520 39.277 1.00 31.14	PROT
	ATOM	137 CD2 TRP 219	29.984 8.585 39.296 1.00 36.40	PROT
	<b>ATOM</b>	138 CE2 TRP 219	30.476 8.547 40.621 1.00 29.24	PROT
	ATOM	139 CE3 TRP 219	30.595 7.781 38.323 1.00 41.61	PROT
30	ATOM	140 CD1 TRP 219	28.771 9.988 40.551 1.00 28.69	PROT
	<b>ATOM</b>	141 NE1 TRP 219	29.718 9.411 41.362 1.00 35.01	PROT
	<b>ATOM</b>	142 CZ2 TRP 219	31.552 7.737 41.004 1.00 30.89	PROT
	ATOM	143 CZ3 TRP 219	31.673 6.969 38.707 1.00 45.72	PROT
	<b>ATOM</b>	144 CH2 TRP 219	32.137 6.958 40.038 1.00 35.17	PROT
35	<b>ATOM</b>	145 C TRP 219	28.125 10.500 35.660 1.00 20.83	PROT
	ATOM	146 O TRP 219	28.467 9.616 34.865 1.00 31.36	PROT
	ATOM	147 N GLU 220	27.143 11.364 35.412 1.00 30.53	PROT
	ATOM	148 CA GLU 220	26.400 11.323 34.159 1.00 33.95	PROT
	ATOM	149 CB GLU 220	25.237 12.318 34.201 1.00 22.17	PROT
40	ATOM	150 C GLU 220	27.356 11.658 33.013 1.00 34.66	PROT
	ATOM	151 O GLU 220	27.233 11.134 31.900 1.00 43.86	PROT
	ATOM	152 N LEU 221	28.320 12.528 33.297 1.00 22.60	PROT
	ATOM	153 CA LEU 221	29.305 12.926 32.304 1.00 17.18	PROT
	ATOM	154 CB LEU 221	29.995 14.219 32.743 1.00 11.03	PROT
45	ATOM	155 CG LEU 221	31.078 14.824 31.850 1.00 5.17	PROT
	ATOM	156 CD1 LEU 221	30.756 14.569 30.415 1.00 6.41	PROT
	ATOM	157 CD2 LEU 221	31.181 16.305 32.092 1.00 10.65	PROT
	ATOM	158 C LEU 221	30.344 11.817 32.122 1.00 22.25	PROT
_	ATOM	159 O LEU 221	30.759 11.521 31.002 1.00 18.99	PROT
50	ATOM	160 N ILE 222	30.754 11.198 33.228 1.00 20.74	PROT

WO 99/26966

Page 322 of 447

	ATOM	161 CA ILE 222 31.744 10.136 33.177 1.00 12.88	PROT
	ATOM	162 CB ILE 222 32.115 9.662 34.587 1.00 12.96	PROT
	ATOM	163 CG2 ILE 222 33.030 8.468 34.515 1.00 2.00	PROT
	ATOM	164 CG1 ILE 222 32.811 10.796 35.332 1.00 16.50	PROT
5	ATOM	165 CD1 ILE 222 33.625 10.351 36.511 1.00 15.90	PROT
	ATOM		PROT
	ATOM		PROT
	ATOM	168 N LYS 223 29.966 8.618 32.530 1.00 33.88	PROT
	ATOM	169 CA LYS 223 29.371 7.503 31.795 1.00 39.02	PROT
10	ATOM	170 CB LYS 223 27.908 7.307 32.224 1.00 40.29	PROT
	ATOM	171 C LYS 223 29.444 7.779 30.293 1.00 39.14	PROT
	ATOM	172 O LYS 223 29.949 6.963 29.517 1.00 32.99	PROT
	ATOM	173 N THR 224 28.936 8.942 29.897 1.00 27.19	PROT
	ATOM	174 CA THR 224 28.929 9.363 28.498 1.00 25.75	PROT
15	ATOM	175 CB THR 224 28.440 10.817 28.407 1.00 22.51	PROT
	ATOM	176 OG1 THR 224 27.018 10.837 28.568 1.00 35.46	PROT
	ATOM	177 CG2 THR 224 28.799 11.436 27.083 1.00 15.53	PROT
	ATOM	178 C THR 224 30.307 9.235 27.833 1.00 22.31	PROT
	ATOM	179 O THR 224 30.480 8.517 26.843 1.00 27.13	PROT
20	ATOM	180 N VAL 225 31.287 9.936 28.386 1.00 17.87	PROT
	ATOM	181 CA VAL 225 32.635 9.906 27.854 1.00 17.07	PROT
	ATOM	182 CB VAL 225 33.559 10.759 28.720 1.00 16.86	PROT
	ATOM	183 CG1 VAL 225 34.845 11.064 27.973 1.00 26.54	
	ATOM	184 CG2 VAL 225 32.854 12.057 29.075 1.00 24.46	
25	ATOM	185 C VAL 225 33.169 8.486 27.793 1.00 16.11	PROT
	ATOM	186 O VAL 225 33.683 8.042 26.763 1.00 12.75	PROT
	ATOM	187 N THR 226 33.040 7.769 28.900 1.00 12.23	PROT
	ATOM	188 CA THR 226 33.520 6.400 28.951 1.00 12.34	PROT
	ATOM	189 CB THR 226 33.175 5.747 30.271 1.00 17.01	PROT
30	ATOM	190 OG1 THR 226 33.715 6.536 31.342 1.00 6.78	PROT
	ATOM	191 CG2 THR 226 33.739 4.324 30.307 1.00 2.00	PROT
	ATOM	192 C THR 226 32.909 5.581 27.837 1.00 14.82	PROT
	ATOM	193 O THR 226 33.623 4.953 27.061 1.00 20.90	PROT
	ATOM	194 N GLU 227 31.582 5.588 27.758 1.00 22.90	PROT
35	ATOM	195 CA GLU 227 30.886 4.849 26.714 1.00 22.63	PROT
	ATOM	196 CB GLU 227 29.417 5.248 26.678 1.00 20.14	PROT
	ATOM	197 C GLU 227 31.556 5.173 25.386 1.00 21.74	PROT
	ATOM	198 O GLU 227 32.057 4.283 24.700 1.00 24.42	PROT
40	ATOM	199 N ALA 228 31.590 6.460 25.050 1.00 13.26	PROT
40	ATOM	200 CA ALA 228 32.196 6.928 23.800 1.00 22.76	PROT
	ATOM	201 CB ALA 228 32.267 8.450 23.785 1.00 22.50	PROT
	ATOM	202 C ALA 228 33.584 6.358 23.538 1.00 19.19	PROT
	ATOM	203 O ALA 228 33.913 6.003 22.408 1.00 17.19	PROT
4.5	ATOM	204 N HIS 229 34.408 6.290 24.573 1.00 20.11	PROT
45	ATOM	205 CA HIS 229 35.741 5.756 24.389 1.00 18.68	PROT
	ATOM	206 CB HIS 229 36.537 5.819 25.686 1.00 10.37	PROT
	ATOM	207 CG HIS 229 37.894 5.201 25.586 1.00 2.00	PROT
	ATOM	208 CD2 HIS 229 38.524 4.299 26.376 1.00 7.61 209 ND1 HIS 229 38.780 5.517 24.582 1.00 3.78	PROT PROT
50	ATOM		PROT
50	ATOM	210 CE1 HIS 229 39.900 4.837 24.758 1.00 15.67	11(01

	ATOM	211 NE2 HIS 229	39.771 4.090 25.840 1.00 7.10	PROT
	ATOM		35.637 4.316 23.940 1.00 21.45	PROT
	ATOM		36.127 3.950 22.866 1.00 22.42	PROT
	ATOM	214 N VAL 230	34.983 3.505 24.762 1.00 21.64	PROT
5	ATOM	215 CA VAL 230	34.827 2.086 24.468 1.00 33.80	PROT
•	ATOM	216 CB VAL 230	33.960 1.388 25.528 1.00 33.11	PROT
	ATOM	217 CG1 VAL 230	34.251 -0.106 25.515 1.00 33.80	PROT
	ATOM	218 CG2 VAL 230	34.228 1.985 26.896 1.00 26.54	PROT
	ATOM	219 C VAL 230	34.224 1.781 23.100 1.00 33.12	PROT
10	ATOM	220 O VAL 230	34.703 0.897 22.385 1.00 40.80	PROT
	ATOM	221 N ALA 231	33.170 2.507 22.746 1.00 36.22	PROT
	ATOM	222 CA ALA 231	32.497 2.298 21.471 1.00 36.24	PROT
	ATOM	223 CB ALA 231	31.318 3.255 21.343 1.00 18.90	PROT
	ATOM	224 C ALA 231	33.445 2.501 20.303 1.00 37.54	PROT
15	ATOM	225 O ALA 231	33.342 1.816 19.285 1.00 35.93	PROT
13	ATOM	226 N THR 232	34.380 3.434 20.474 1.00 23.74	PROT
	ATOM	227 CA THR 232	35.329 3.789 19.432 1.00 15.54	PROT
	ATOM	228 CB THR 232	35.335 5.321 19.238 1.00 9.70	PROT
	ATOM	229 OG1 THR 232	35.733 5.949 20.460 1.00 16.73	PROT
20	ATOM	230 CG2 THR 232	33.942 5.828 18.891 1.00 2.00	PROT
20	ATOM	231 C THR 232	36.758 3.309 19.670 1.00 19.86	PROT
	ATOM	232 O THR 232	37.695 3.854 19.094 1.00 15.31	PROT
	ATOM	233 N ASN 233	36.938 2.305 20.523 1.00 28.26	PROT
	ATOM	234 CA ASN 233	38.280 1.771 20.772 1.00 39.32	PROT
25	ATOM	235 CB ASN 233	38.435 1.343 22.234 1.00 47.14	PROT
	ATOM	236 CG ASN 233	39.804 1.689 22.801 1.00 54.02	PROT
	ATOM	237 OD1 ASN 233	40.633 2.303 22.128 1.00 60.36	PROT
	ATOM	238 ND2 ASN 233	40.045 1.296 24.045 1.00 48.67	PROT
	ATOM	239 C ASN 233	38.507 0.574 19.840 1.00 49.33	PROT
30	ATOM	240 O ASN 233	38.338 0.693 18.625 1.00 65.36	PROT
	ATOM	241 N ALA 234	38.877 -0.577 20.388 1.00 57.89	PROT
	ATOM	242 CA ALA 234	39.090 -1.752 19.552 1.00 57.22	PROT
	ATOM	243 CB ALA 234	40.372 -1.595 18.754 1.00 48.03	PROT
	ATOM	244 C ALA 234	39.141 -3.027 20.384 1.00 62.42	PROT
35	ATOM	245 O ALA 234	38.471 -3.073 21.440 1.00 56.93	PROT
	ATOM	246 OT ALA 234	39.853 -3.968 19.965 1.00 76.16	PROT
	ATOM	247 N TRP 239	41.987 -7.449 22.970 1.00 58.82	PROT
	ATOM	248 CA TRP 239	43.077 -6.886 22.154 1.00 51.37	PROT
	ATOM	249 CB TRP 239	43.325 -5.406 22.534 1.00 45.12	PROT
40	ATOM	250 CG TRP 239	44.193 -5.170 23.760 1.00 43.09	PROT
	ATOM	251 CD2 TRP 239	45.617 -5.037 23.793 1.00 32.36	PROT
	ATOM	252 CE2 TRP 239	45.990 -4.872 25.142 1.00 28.37	PROT
	ATOM	253 CE3 TRP 239	46.615 -5.049 22.813 1.00 40.79	PROT
	ATOM	254 CD1 TRP 239	43.773 -5.073 25.059 1.00 46.63	PROT
45	ATOM	255 NE1 TRP 239	44.847 -4.896 25.893 1.00 27.08	PROT
•	ATOM	256 CZ2 TRP 239	47.315 -4.717 25.535 1.00 35.48	PROT.
	ATOM	257 CZ3 TRP 239	47.936 -4.896 23.204 1.00 40.18	PROT
	ATOM	258 CH2 TRP 239	48.273 -4.733 24.554 1.00 49.93	PROT
	ATOM	259 C TRP 239	44.422 -7.623 22.063 1.00 49.76	PROT
50	ATOM	260 O TRP 239	44.944 -7.799 20.962 1.00 48.14	PROT

	<b>ATOM</b>	261 N LYS 240	44.975 -8.048 23.198 1.00 38.92	PROT
	<b>ATOM</b>	262 CA LYS 240	46.263 -8.735 23.232 1.00 37.29	PROT
	<b>ATOM</b>	263 CB LYS 240	46.572 -9.196 24.657 1.00 38.79	PROT
	<b>ATOM</b>	264 CG LYS 240	47.106 -8.099 25.571 1.00 38.43	PROT
5	<b>ATOM</b>	265 CD LYS 240	48.307 -8.584 26.370 1.00 35.71	PROT
	<b>ATOM</b>	266 CE LYS 240	48.631 -7.646 27.523 1.00 37.87	PROT
	ATOM	267 NZ LYS 240	49.058 -8.377 28.750 1.00 28.85	PROT
	<b>ATOM</b>	268 C LYS 240	46.404 -9.914 22.269 1.00 42.18	PROT
	<b>ATOM</b>	269 O LYS 240	47.491 -10.132 21.732 1.00 45.89	PROT
10	ATOM	270 N GLN 241	45.331 -10.679 22.058 1.00 46.08	PROT
	ATOM	271 CA GLN 241	45.390 -11.816 21.133 1.00 45.02	PROT
	ATOM	272 CB GLN 241	44.575 -13.011 21.638 1.00 46.30	PROT
	ATOM	273 CG GLN 241	44.284 -13.018 23.116 1.00 60.38	PROT
	<b>ATOM</b>	274 CD GLN 241	42.828 -13.312 23.408 1.00 63.76	PROT
15	ATOM	275 OE1 GLN 241	42.154 -13.988 22.631 1.00 66.34	PROT
	ATOM	276 NE2 GLN 241	42.333 -12.801 24.531 1.00 69.18	PROT
	<b>ATOM</b>	277 C GLN 241	44.866 -11.405 19.764 1.00 45.77	PROT
	ATOM	278 O GLN 241	45.107 -12.085 18.765 1.00 51.18	PROT
	ATOM	279 N LYS 242	44.132 -10.300 19.723 1.00 42.04	PROT
20	ATOM	280 CA LYS 242	43.613 -9.794 18.464 1.00 48.33	PROT
	ATOM	281 CB LYS 242	42.498 -8.786 18.727 1.00 40.17	PROT
	ATOM	282 C LYS 242	44.796 -9.123 17.742 1.00 53.04	PROT
	ATOM	283 O LYS 242	44.709 -8.753 16.565 1.00 48.21	PROT
	ATOM	284 N ARG 243	45.906 -8.992 18.470 1.00 45.44	PROT
25	ATOM	285 CA ARG 243	47.128 -8.374 17.965 1.00 43.53	PROT
	ATOM	286 CB ARG 243	48.108 -8.135 19.118 1.00 40.21	PROT
	ATOM	287 C ARG 243	47.795 -9.220 16.892 1.00 45.96	PROT
	ATOM	288 O ARG 243	47.684 -10.443 16.894 1.00 50.22	PROT
	ATOM	289 N LYS 244	48.498 -8.551 15.982 1.00 52.12	PROT
30	ATOM	290 CA LYS 244	49.202 -9.202 14.879 1.00 45.30	PROT
	ATOM	291 CB LYS 244	48.466 -8.950 13.558 1.00 48.24	PROT
	ATOM	292 CG LYS 244	47.109 -9.631 13.446 1.00 53.78	PROT
	ATOM	293 CD LYS 244	46.835 -10.078 12.011 1.00 60.50	PROT
	ATOM	294 CE LYS 244	46.038 -9.030 11.241 1.00 61.03	PROT
35	ATOM	295 NZ LYS 244	45.455 -7.997 12.146 1.00 55.25	PROT
	ATOM	296 C LYS 244	50.616 -8.641 14.786 1.00 40.33	PROT
	ATOM	297 O LYS 244	50.849 -7.629 14.125 1.00 36.07	PROT
	ATOM	298 N PHE 245	51.556 -9.312 15.445 1.00 27.87	PROT
	ATOM	299 CA PHE 245	52.949 -8.885 15.461 1.00 30.61	PROT
40	ATOM	300 CB PHE 245	53.784 -9.887 16.253 1.00 20.28	PROT
	ATOM	301 CG PHE 245	53.454 -9.922 17.713 1.00 37.23	PROT
	ATOM	302 CD1 PHE 245	52.636 -10.917 18.234 1.00 40.93	PROT
	ATOM	303 CD2 PHE 245	53.958 -8.959 18.577 1.00 41.60	PROT
	ATOM	304 CE1 PHE 245	52.326 -10.953 19.594 1.00 42.54	PROT
45	ATOM	305 CE2 PHE 245	53.652 -8.989 19.936 1.00 45.84	PROT
	ATOM	306 CZ PHE 245	52.835 -9.988 20.443 1.00 33.72	PROT
	ATOM	307 C PHE 245	53.549 -8.693 14.068 1.00 38.75	PROT
	ATOM	308 O PHE 245	53.794 -9.660 13.337 1.00 48.93	PROT
	ATOM	309 N LEU 246	53.789 -7.437 13.704 1.00 41.18	PROT
50	ATOM	310 CA LEU 246	54.362 -7.124 12.404 1.00 43.43	PROT
	· <del>-</del> - · -			

	4 TO 1 6	211 CD TELL 246	54 279 5 612 12 101 1 00 42 79	ррот
	ATOM		54.378 -5.612 12.181 1.00 42.78	PROT PROT
	ATOM	312 CG LEU 246	54.535 -5.200 10.718 1.00 49.88 53.528 -4.113 10.365 1.00 40.64	PROT
	ATOM	313 CD1 LEU 246	55.966 -4.730 10.485 1.00 48.66	
_	ATOM	314 CD2 LEU 246	55.777 -7.692 12.250 1.00 42.60	PROT
5	ATOM		56.677 -7.383 13.028 1.00 45.75	PROT
	ATOM		55,977 -8.540 11.233 1.00 50.03	PROT
	ATOM		54.914 -8.924 10.286 1.00 50.03	PROT
	ATOM	318 CD PRO 247	57.237 -9.199 10.894 1.00 49.90	PROT
10	ATOM	319 CA PRO 247	57.181 -9.282 9.369 1.00 59.51	PROT
10	ATOM	320 CB PRO 247	55.678 -9.244 9.023 1.00 52.86	PROT
	ATOM	321 CG PRO 247		
	ATOM		58.499 -8.494 11.392 1.00 48.85	PROT
	ATOM		58.675 -7.295 11.186 1.00 49.28	PROT
	ATOM		59.379 -9.261 12.032 1.00 47.62	PROT
15	ATOM	325 CA GLU 248	60.628 -8.733 12.574 1.00 51.41	PROT
	ATOM	326 CB GLU 248	61.266 -9.750 13.522 1.00 44.22	PROT
	ATOM		51.623 -8.354 11.490 1.00 53.28	PROT
	ATOM		52.815 -8.214 11.765 1.00 62.57	PROT
••	ATOM		51.146 -8.200 10.258 1.00 56.20	PROT
20	ATOM		62.030 -7.818 9.164 1.00 55.88	PROT
	ATOM		62.231 -8.981 8.173 1.00 53.88	PROT
	ATOM		60.928 -9.637 7.739 1.00 54.39	PROT
	ATOM	333 OD1 ASP 249	60.578 -10.693 8.310 1.00 57.70	PROT
0.5	ATOM	334 OD2 ASP 249	60.264 -9.112 6.819 1.00 45.76	PROT
25	ATOM	<del>-</del>	51.539 -6.567 8.437 1.00 54.20	PROT
	ATOM		52.119 -6.154 7.429 1.00 55.31	PROT
	ATOM		0.469 -5.965 8.954 1.00 46.13	PROT
	ATOM		59.933 -4.735 8.376 1.00 46.12	PROT
	ATOM		58.413 -4.764 8.253 1.00 43.38	PROT
30	ATOM		57.892 -3.344 8.057 1.00 39.15	PROT
	ATOM		58.007 -5.654 7.074 1.00 48.96	PROT
	ATOM		56.707 -6.401 7.283 1.00 43.14	PROT
	ATOM		0.311 -3.590 9.294 1.00 45.32	PROT
	ATOM		0.257 -3.724 10.513 1.00 43.74	PROT
35	ATOM		60.680 -2.459 8.711 1.00 36.80	PROT
	ATOM	346 CA GLY 251	61.091 -1.329 9.521 1.00 39.28	PROT
	ATOM	-	52.370 -1.621 10.305 1.00 44.31	PROT
	ATOM		62.538 -1.145 11.428 1.00 51.39	PROT
	ATOM		63.277 -2.399 9.715 1.00 55.47	PROT
40	ATOM	350 CA GLN 252	64.536 -2.745 10.374 1.00 54.24	PROT
	ATOM	351 CB GLN 252	64.792 -4.237 10.245 1.00 49.31	PROT
	ATOM		55.720 -1.959 9.812 1.00 54.86	PROT
	ATOM		65.492 -1.079 8.953 1.00 58.80	PROT
	ATOM	354 CB VAL 264	60.887 6.759 5.510 1.00 34.33	PROT
45	ATOM	355 CG1 VAL 264	59.550 6.086 5.790 1.00 34.34	PROT
	ATOM	356 CG2 VAL 264	60.893 8.163 6.080 1.00 20.22	PROT
	ATOM		62.053 4.557 5.439 1.00 34.08	PROT
	ATOM		62.280 4.466 4.232 1.00 46.39	PROT
	ATOM		63.361 6.605 5.966 1.00 21.27	PROT
50	ATOM	360 CA VAL 264	62.041 5.920 6.122 1.00 29.68	PROT

	ATOM	361 N ASP 265	61.809 3.499 6.209 1.00 40.63	PROT
	ATOM	362 CA ASP 265	61.796 2.141 5.670 1.00 43.58	PROT
	ATOM	363 CB ASP 265	61.243 1.160 6.704 1.00 44.07	PROT
	ATOM	364 CG ASP 265	61.179 -0.262 6.185 1.00 49.19	PROT
5	ATOM	365 OD1 ASP 265	62.223 -0.945 6.175 1.00 57.67	PROT
	ATOM	366 OD2 ASP 265	60.082 -0.702 5.789 1.00 54.75	PROT
	ATOM	367 C ASP 265	60.956 2.071 4.401 1.00 48.03	PROT
	ATOM	368 O ASP 265	61.362 1.458 3.411 1.00 57.44	PROT
	ATOM	369 N LEU 266	59.793 2.711 4.436 1.00 40.55	PROT
10	ATOM	370 CA LEU 266	58.879 2.741 3.295 1.00 45.78	PROT
10	ATOM	371 CB LEU 266	59.638 2.962 1.977 1.00 45.92	PROT
	ATOM	372 CG LEU 266	59.881 4.407 1.506 1.00 48.41	PROT
	ATOM	373 CDI LEU 266	59.934 4.432 -0.007 1.00 32.83	PROT
	ATOM	374 CD2 LEU 266	58.787 5.344 2.012 1.00 45.08	PROT
15	ATOM	375 C LEU 266	58.064 1.462 3.214 1.00 45.45	PROT
	ATOM	376 O LEU 266	56.862 1.503 2.949 1.00 42.92	PROT
	ATOM	377 N GLU 267	58.712 0.324 3.431 1.00 46.47	PROT
	ATOM	378 CA GLU 267	57.986 -0.935 3.415 1.00 44.34	PROT
	ATOM	379 CB GLU 267	58.943 -2.123 3.505 1.00 39.42	PROT
20	ATOM	380 CG GLU 267	58.291 -3.457 3.188 1.00 40.68	PROT
	ATOM	381 CD GLU 267	58.929 -4.607 3.943 1.00 63.54	PROT
	ATOM	382 OE1 GLU 267	60.103 -4.470 4.361 1.00 68.92	PROT
	<b>ATOM</b>	383 OE2 GLU 267	58.258 -5.650 4.120 1.00 66.66	PROT
	<b>ATOM</b>	384 C GLU 267	57.106 -0.880 4.655 1.00 41.57	PROT
25	<b>ATOM</b>	385 O GLU 267	55.991 -1.398 4.673 1.00 48.68	PROT
	<b>ATOM</b>	386 N ALA 268	57.620 -0.215 5.686 1.00 39.33	PROT
	<b>ATOM</b>	387 CA ALA 268	56.916 -0.057 6.951 1.00 31.62	PROT
	ATOM	388 CB ALA 268	57.918 0.134 8.063 1.00 7.56	PROT
	<b>ATOM</b>	389 C ALA 268	55.960 1.135 6.888 1.00 25.96	PROT
30	ATOM	390 O ALA 268	54.786 1.036 7.237 1.00 17.35	PROT
	ATOM	391 N PHE 269	56.464 2.274 6.446 1.00 11.34	PROT
	ATOM	392 CA PHE 269	55.615 3.453 6.335 1.00 15.72	PROT
	ATOM	393 CB PHE 269	56.274 4.474 5.405 1.00 20.08	PROT
	ATOM	394 CG PHE 269	55.552 5.788 5.334 1.00 24.67	PROT
35	ATOM	395 CD1 PHE 269	55.661 6.713 6.369 1.00 15.69	PROT
	ATOM	396 CD2 PHE 269	54.772 6.111 4.222 1.00 20.64	PROT
	ATOM	397 CE1 PHE 269	55.003 7.942 6.300 1.00 22.55	PROT
	ATOM	398 CE2 PHE 269	54.108 7.342 4.143 1.00 19.77	PROT
	ATOM	399 CZ PHE 269	54.224 8.257 5.186 1.00 19.27	PROT
40	ATOM	400 C PHE 269	54.277 3.010 5.754 1.00 19.45	PROT
	ATOM	401 O PHE 269	53.212 3.351 6.261 1.00 13.40	PROT
	ATOM	402 N SER 270	54.367 2.214 4.692 1.00 43.85	PROT
	ATOM	403 CA SER 270	53.217 1.686 3.967 1.00 46.67	PROT
	ATOM	404 CB SER 270	53.687 0.669 2.924 1.00 53.60	PROT
45	ATOM	405 OG SER 270	52.662 0.382 1.988 1.00 68.82	PROT
	ATOM	406 C SER 270	52.181 1.039 4.865 1.00 43.32	PROT
	ATOM	407 O SER 270	51.024 1.459 4.893 1.00 43.87	PROT
	ATOM	408 N HIS 271	52.594 0.009 5.590 1.00 34.59	PROT
50	ATOM	409 CA HIS 271	51.681 -0.694 6.486 1.00 37.12	PROT
50	ATOM	410 CB HIS 271	52.441 -1.772 7.266 1.00 46.61	PROT

	ATOM	411 CG HIS 271 52.603 -3.056 6.512 1.00 63.99	PROT
	ATOM	412 CD2 HIS 271 51.879 -4.201 6.533 1.00 62.06	PROT
	ATOM	413 ND1 HIS 271 53.608 -3.256 5.590 1.00 60.86	PROT
	ATOM	414 CE1 HIS 271 53.497 -4.467 5.075 1.00 60.70	PROT
5	ATOM	415 NE2 HIS 271 52.456 -5.061 5.630 1.00 64.10	<b>PROT</b>
-	ATOM		PROT
	ATOM		PROT
	ATOM		PROT
	ATOM	419 CA PHE 272 51.190 2.038 9.085 1.00 27.77	PROT
10	ATOM	420 CB PHE 272 52.302 2.886 9.714 1.00 10.49	PROT
10	ATOM	421 CG PHE 272 53.338 2.086 10.459 1.00 6.98	PROT
	ATOM	422 CD1 PHE 272 54.671 2.478 10.449 1.00 4.13	PROT
		423 CD2 PHE 272 52.978 0.961 11.193 1.00 6.95	PROT
	ATOM		PROT
1.5	ATOM		PROT
15	ATOM	425 CE2 PHE 272 53.930 0.242 11.909 1.00 6.13	
	ATOM	426 CZ PHE 272 55.263 0.645 11.895 1.00 8.93	PROT
	ATOM		PROT
	ATOM		PROT
	ATOM	1	PROT
20	ATOM	430 CA THR 273 49.633 4.343 6.487 1.00 33.39	PROT
	ATOM	431 CB THR 273 50.335 4.912 5.243 1.00 36.80	PROT
	ATOM	432 OG1 THR 273 50.649 3.847 4.332 1.00 27.42	PROT
	ATOM	433 CG2 THR 273 51.613 5.641 5.656 1.00 32.25	PROT
	ATOM		PROT
25	ATOM		PROT
	ATOM	· · · · · · · · · · · · · · · · · · ·	PROT
	ATOM	437 CA LYS 274 47.196 1.555 5.726 1.00 42.17	PROT
	ATOM	438 CB LYS 274 47.544 0.069 5.615 1.00 40.02	PROT
	ATOM	439 C LYS 274 46.153 1.778 6.818 1.00 41.47	PROT
30	ATOM		PROT
	ATOM	441 N ILE 275 46.456 1.290 8.019 1.00 34.08 F	PROT
	ATOM	442 CA ILE 275 45.559 1.403 9.166 1.00 25.49	PROT
	ATOM	443 CB ILE 275 45.991 0.435 10.262 1.00 19.72	PROT
	ATOM	444 CG2 ILE 275 46.290 -0.934 9.642 1.00 23.39	<b>PROT</b>
35	<b>ATOM</b>	445 CG1 ILE 275 47.249 0.958 10.953 1.00 12.96	<b>PROT</b>
	<b>ATOM</b>	446 CD1 ILE 275 47.970 -0.103 11.769 1.00 11.07	PROT
	ATOM		PROT
	ATOM		PROT
	ATOM		ROT
40	ATOM		PROT
	ATOM		PROT
	ATOM	452 CG2 ILE 276 46.894 6.267 7.831 1.00 27.28	PROT
	ATOM	453 CG1 ILE 276 47.374 7.288 10.028 1.00 6.75	PROT
	ATOM	454 CD1 ILE 276 48.349 7.255 11.153 1.00 15.44	PROT
45	ATOM		PROT
43	ATOM		PROT
			PROT
	ATOM	457 N THR 277 44.303 3.411 8.333 1.00 22.12 458 CA THR 277 43.007 6.005 8.260 1.00 27.16	PROT
	ATOM		PROT
50	ATOM		PROT
50	ATOM	460 OG1 THR 277 43.665 5.584 5.955 1.00 22.55	LVOI

	ATOM	461 CG2 THR 277 41.594 6.763 6.337 1.00 26.98	PROT
	ATOM	462 C THR 277 41.944 5.591 9.270 1.00 25.23	PROT
	ATOM	463 O THR 277 41.271 6.443 9.847 1.00 21.62	PROT
	ATOM	464 N PRO 278 41.769 4.279 9.491 1.00 18.64	PROT
5	ATOM	465 CD PRO 278 42.472 3.167 8.832 1.00 9.52	PROT
	ATOM	466 CA PRO 278 40.765 3.803 10.453 1.00 18.48	PROT
	<b>ATOM</b>	467 CB PRO 278 40.907 2.280 10.415 1.00 14.77	PROT
	ATOM	468 CG PRO 278 42.195 2.008 9.738 1.00 7.70	PROT
	<b>ATOM</b>	469 C PRO 278 40.956 4.356 11.870 1.00 25.40	PROT
10	<b>ATOM</b>	470 O PRO 278 39.983 4.628 12.576 1.00 22.33	PROT
	<b>ATOM</b>	471 N ALA 279 42.211 4.507 12.285 1.00 22.14	PROT
	<b>ATOM</b>	472 CA ALA 279 42.519 5.038 13.607 1.00 20.26	PROT
	<b>ATOM</b>	473 CB ALA 279 44.016 5.033 13.831 1.00 13.33	PROT
	ATOM	474 C ALA 279 41.984 6.456 13.699 1.00 16.49	PROT
15	ATOM	475 O ALA 279 41.222 6.797 14.598 1.00 32.38	PROT
	ATOM	476 N ILE 280 42.384 7.286 12.753 1.00 7.56 H	PROT
	ATOM	477 CA ILE 280 41.935 8.666 12.734 1.00 9.96	PROT
	ATOM		PROT
	ATOM	479 CG2 ILE 280 42.172 10.871 11.581 1.00 2.00	PROT
20	ATOM	480 CG1 ILE 280 43.901 9.059 11.220 1.00 10.96	<b>PROT</b>
	ATOM	481 CD1 ILE 280 44.615 10.036 10.294 1.00 8.54	PROT
	ATOM		PROT
	ATOM		PROT
	<b>ATOM</b>	484 N THR 281 39.692 7.883 12.172 1.00 24.18	PROT
25	<b>ATOM</b>	485 CA THR 281 38.238 7.962 12.153 1.00 24.77	PROT
	ATOM	486 CB THR 281 37.650 6.952 11.145 1.00 33.90	PROT
	ATOM	487 OG1 THR 281 38.607 6.711 10.108 1.00 34.62	PROT
	ATOM	488 CG2 THR 281 36.379 7.506 10.513 1.00 39.80	PROT
	<b>ATOM</b>	489 C THR 281 37.655 7.726 13.535 1.00 23.39	PROT
30	ATOM	490 O THR 281 36.733 8.422 13.960 1.00 19.51	PROT
	ATOM	491 N ARG 282 38.213 6.743 14.234 1.00 16.90	PROT
	ATOM	492 CA ARG 282 37.781 6.404 15.583 1.00 12.29	PROT
	ATOM	493 CB ARG 282 38.641 5.260 16.115 1.00 5.36	PROT
	ATOM	494 CG ARG 282 37.936 3.926 16.136 1.00 17.05	PROT
35	ATOM	495 CD ARG 282 38.296 3.095 14.942 1.00 18.41	PROT
	ATOM	496 NE ARG 282 39.622 2.475 15.011 1.00 35.77	PROT
	<b>ATOM</b>	497 CZ ARG 282 40.454 2.501 16.055 1.00 36.80	PROT
	ATOM	498 NH1 ARG 282 41.629 1.888 15.967 1.00 35.96	PROT
	ATOM	499 NH2 ARG 282 40.134 3.120 17.183 1.00 25.20	PROT
40	ATOM	500 C ARG 282 37.863 7.626 16.520 1.00 16.75	PROT
	ATOM	501 O ARG 282 37.078 7.758 17.456 1.00 22.98	PROT
	ATOM	502 N VAL 283 38.813 8.518 16.268 1.00 11.92	PROT
	ATOM	503 CA VAL 283 38.937 9.719 17.083 1.00 14.68	PROT
	<b>ATOM</b>	504 CB VAL 283 40.191 10.541 16.696 1.00 23.35	PROT
45	ATOM	505 CG1 VAL 283 40.467 11.593 17.752 1.00 11.98	PROT
	ATOM	506 CG2 VAL 283 41.396 9.621 16.526 1.00 20.41	PROT
	ATOM	507 C VAL 283 37.705 10.580 16.833 1.00 12.72	PROT
	ATOM	508 O VAL 283 36.965 10.929 17.752 1.00 20.37	PROT
	ATOM	509 N VAL 284 37.503 10.920 15.567 1.00 18.28	<b>PROT</b>
50	ATOM	510 CA VAL 284 36.369 11.727 15.150 1.00 16.98	PROT

	ATOM	511 CB VAL 284	36.251 11.765 13.602 1.00 27.40	PROT
	ATOM	512 CG1 VAL 284	35.434 12.973 13.172 1.00 19.30 37.649 11.794 12.959 1.00 16.94	
	ATOM	513 CG2 VAL 284		
_	ATOM	514 C VAL 284	35.113 11.093 15.715 1.00 14.89	PROT PROT
5	ATOM	515 O VAL 284	34.233 11.781 16.219 1.00 10.93 35.046 9.768 15.623 1.00 10.68	
	ATOM	516 N ASP 285		PROT
	ATOM	517 CA ASP 285		PROT
	ATOM	518 CB ASP 285	34.079 7.518 15.874 1.00 22.99 33.985 7.130 14.397 1.00 30.01	PROT
10	ATOM	519 CG ASP 285 520 OD1 ASP 285		PROT PROT
10	ATOM			
	ATOM	521 OD2 ASP 285	34.720 6.202 13.993 1.00 27.74 33.734 9.274 17.604 1.00 26.87	PROT PROT
	ATOM	522 C ASP 285		PROT
	ATOM	523 O ASP 285	32.609 9.349 18.103 1.00 39.89 34.861 9.405 18.308 1.00 25.45	PROT
1.5	ATOM	524 N PHE 286 525 CA PHE 286		
15	ATOM			PROT
	ATOM	526 CB PHE 286 527 CG PHE 286	36.284 9.533 20.305 1.00 7.30 36.454 10.104 21.703 1.00 17.92	PROT
	ATOM ATOM	527 CG PHE 286 528 CD1 PHE 286	35.848 9.499 22.805 1.00 17.92	PROT PROT
		529 CD2 PHE 286	37.229 11.245 21.920 1.00 19.33	
20	ATOM ATOM	530 CE1 PHE 286	36.014 10.021 24.087 1.00 19.24	PROT PROT
20	ATOM	530 CE1 PHE 286	37.395 11.769 23.207 1.00 11.33	PROT
	ATOM	531 CE2 PHE 286	36.786 11.154 24.283 1.00 2.00	PROT
	ATOM	532 CZ FHE 286	34.313 11.043 20.030 1.00 17.67	PROT
	ATOM	534 O PHE 286	33.367 11.201 20.797 1.00 14.36	PROT
25	ATOM	534 O FHE 280 535 N ALA 287	34.905 12.056 19.410 1.00 12.57	PROT
23	ATOM	536 CA ALA 287	34.443 13.426 19.622 1.00 12.49	PROT
	ATOM	537 CB ALA 287	35.250 14.386 18.759 1.00 23.54	PROT
	ATOM	538 C ALA 287	32.954 13.559 19.307 1.00 9.21	PROT
	ATOM	539 O ALA 287	32.209 14.205 20.043 1.00 11.68	PROT
30	ATOM	540 N LYS 288	32.540 12.929 18.209 1.00 16.43	PROT
50	ATOM	541 CA LYS 288	31.157 12.944 17.736 1.00 16.10	PROT
	ATOM	542 CB LYS 288	31.003 11.977 16.569 1.00 13.15	PROT
	ATOM	543 CG LYS 288	31.117 12.636 15.219 1.00 25.55	PROT
	ATOM	544 CD LYS 288	30.480 11.779 14.136 1.00 32.95	PROT
35	ATOM	545 CE LYS 288	31.279 10.507 13.900 1.00 34.58	PROT
	ATOM	546 NZ LYS 288	30.755 9.721 12.748 1.00 36.93	PROT
	ATOM	547 C LYS 288	30.154 12.569 18.813 1.00 18.87	PROT
	ATOM	548 O LYS 288	29.078 13.171 18.917 1.00 12.83	PROT
	ATOM	549 N LYS 289	30.525 11.574 19.614 1.00 11.81	PROT
40	<b>ATOM</b>	550 CA LYS 289	29.674 11.067 20.681 1.00 15.53	PROT
	<b>ATOM</b>	551 CB LYS 289	30.070 9.631 21.011 1.00 15.88	PROT
	<b>ATOM</b>	552 CG LYS 289	29.767 8.645 19.911 1.00 20.93	PROT
	<b>ATOM</b>	553 CD LYS 289	29.140 7.382 20.471 1.00 28.97	PROT
	<b>ATOM</b>	554 CE LYS 289	29.951 6.167 20.071 1.00 25.06	PROT
45	<b>ATOM</b>	555 NZ LYS 289	30.043 6.060 18.590 1.00 39.19	PROT
	<b>ATOM</b>	556 C LYS 289	29.660 11.884 21.969 1.00 15.95	PROT
	ATOM	557 O LYS 289	29.205 11.398 23.001 1.00 28.53	PROT
	ATOM	558 N LEU 290	30.151 13.116 21.919 1.00 10.13	PROT
	ATOM	559 CA LEU 290	30.155 13.959 23.104 1.00 7.83	PROT
50	ATOM	560 CB LEU 290	31.588 14.300 23.532 1.00 14.46	PROT

	ATOM	561 CG LEU 290	32.676 13.228 23.542 1.00 11.22	PROT
	ATOM	562 CD1 LEU 290	34.016 13.900 23.678 1.00 3.02	PROT
	<b>ATOM</b>	563 CD2 LEU 290	32.449 12.257 24.686 1.00 9.39	PROT
	ATOM	564 C LEU 290	29.410 15.259 22.849 1.00 7.59	PROT
5	ATOM	565 O LEU 290	29.942 16.148 22.196 1.00 11.01	PROT
	ATOM	566 N PRO 291	28.169 15.381 23.365 1.00 14.33	PROT
	ATOM	567 CD PRO 291	27.515 14.291 24.109 1.00 18.52	PROT
	ATOM	568 CA PRO 291	27.290 16.556 23.240 1.00 6.61	PROT
	ATOM	569 CB PRO 291	26.296 16.400 24.384 1.00 11.95	PROT
10	ATOM	570 CG PRO 291	26,496 15.004 24.929 1.00 20.22	PROT
	<b>ATOM</b>	571 C PRO 291	28.029 17.885 23.332 1.00 14.74	PROT
	<b>ATOM</b>	572 O PRO 291	27.795 18.792 22.537 1.00 26.09	PROT
	<b>ATOM</b>	573 N MET 292	28.917 18.002 24.315 1.00 24.06	PROT
	<b>ATOM</b>	574 CA MET 292	29.697 19.225 24.494 1.00 25.33	PROT
15	ATOM	575 CB MET 292	30.706 19.046 25.628 1.00 26.65	PROT
	<b>ATOM</b>	576 CG MET 292	30.222 19.581 26.962 1.00 26.97	PROT
	ATOM	577 SD MET 292	31.153 18.943 28.362 1.00 29.01	PROT
	ATOM	578 CE MET 292	30.315 17.438 28.685 1.00 17.91	PROT
	ATOM	579 C MET 292	30.430 19.588 23.204 1.00 23.01	PROT
20	ATOM	580 O MET 292	30.478 20.747 22.813 1.00 31.98	PROT
	ATOM	581 N PHE 293	31.007 18.591 22.547 1.00 23.44	PROT
	ATOM	582 CA PHE 293	31.724 18.819 21.297 1.00 24.83	PROT
	ATOM	583 CB PHE 293	32.389 17.529 20.830 1.00 15.05	PROT
	ATOM	584 CG PHE 293	33.214 17.686 19.594 1.00 13.55	PROT
25	ATOM	585 CD1 PHE 293	34.376 18.446 19.614 1.00 19.86	PROT
	ATOM	586 CD2 PHE 293	32.867 17.024 18.425 1.00 22.99	PROT
	ATOM	587 CE1 PHE 293	35.184 18.540 18.495 1.00 18.15	PROT
	ATOM	588 CE2 PHE 293	33.671 17.108 17.291 1.00 20.83	PROT
	ATOM	589 CZ PHE 293	34.831 17.866 17.328 1.00 22.53	PROT
30	ATOM	590 C PHE 293	30.759 19.291 20.222 1.00 27.26	PROT
	ATOM	591 O PHE 293	30.971 20.319 19.577 1.00 28.69	PROT
	ATOM	592 N CYS 294	29.689 18.528 20.040 1.00 29.92	PROT
	ATOM	593 CA CYS 294	28.700 18.855 19.037 1.00 35.54	PROT
	ATOM	594 CB CYS 294	27.540 17.860 19.106 1.00 19.11	PROT
35	ATOM	595 SG CYS 294	27.843 16.358 18.132 1.00 35.66	PROT
	ATOM	596 C CYS 294	28.203 20.291 19.171 1.00 38.84	PROT
	ATOM	597 O CYS 294	28.072 20.995 18.169 1.00 45.94	PROT
	ATOM	598 N GLU 295 599 CA GLU 295	27.959 20.739 20.401 1.00 27.34 27.472 22.097 20.632 1.00 21.06	PROT
40	ATOM			PROT PROT
40	ATOM	600 CB GLU 295	27.178 22.306 22.121 1.00 29.78	PROT
	ATOM	601 C GLU 295	28.458 23.158 20.128 1.00 23.67 28.228 24.357 20.272 1.00 29.89	PROT
	ATOM ATOM	602 O GLU 295 603 N LEU 296	29.551 22.715 19.522 1.00 29.89	PROT
	ATOM	603 N LEU 296 604 CA LEU 296	30.545 23.642 19.005 1.00 26.35	PROT
45	ATOM	605 CB LEU 296	31.947 23.128 19.330 1.00 25.17	PROT
4)	ATOM	606 CG LEU 296	32.419 23.157 20.778 1.00 23.17	PROT
	ATOM	607 CD1 LEU 296	33.593 22.217 20.931 1.00 23.61	PROT
	ATOM	607 CD1 LEU 296	32.814 24.564 21.160 1.00 13.82	PROT
	ATOM	609 C LEU 296	30.415 23.783 17.493 1.00 31.88	PROT
50	ATOM	610 O LEU 296	29.890 22.890 16.827 1.00 45.99	PROT
50	111 0111	010 O EEO 270	27.070 22.070 10.027 1.00 13.77	1101

	ATOM	611 N PRO 297	30.884 24.912 16.932 1.00 27.00	PROT
	ATOM	612 CD PRO 297	31.423 26.037 17.708 1.00 36.12	PROT
	ATOM	613 CA PRO 297	30.856 25.222 15.492 1.00 22.30	PROT
	ATOM	614 CB PRO 297	31.182 26.716 15.424 1.00 16.06	PROT
5	ATOM	615 CG PRO 297	31.107 27.208 16.827 1.00 42.41	PROT
•	ATOM	616 C PRO 297	31.838 24.413 14.642 1.00 28.19	PROT
	ATOM	617 O PRO 297	32.983 24.189 15.036 1.00 39.38	PROT
	ATOM	618 N CYS 298	31.371 24.014 13.457 1.00 35.37	PROT
	ATOM	619 CA CYS 298	32.134 23.233 12.481 1.00 32.41	PROT
10	ATOM	620 CB CYS 298	31.416 23.289 11.112 1.00 40.85	PROT
	ATOM	621 SG CYS 298	32.431 23.615 9.614 1.00 61.24	PROT
	ATOM	622 C CYS 298	33,596 23.654 12.352 1.00 31.68	PROT
	ATOM	623 O CYS 298	34.474 22.804 12.225 1.00 28.49	PROT
	ATOM	624 N GLU 299	33.869 24.954 12.393 1.00 29.93	PROT
15	ATOM	625 CA GLU 299	35.253 25.407 12.278 1.00 36.38	PROT
• •	ATOM	626 CB GLU 299	35.346 26.931 12.203 1.00 32.78	PROT
	ATOM	627 CG GLU 299	34.467 27.546 11.167 1.00 43.40	PROT
	ATOM	628 CD GLU 299	33.038 27.593 11.625 1.00 58.19	PROT
	ATOM	629 OE1 GLU 299		PROT
20	ATOM	630 OE2 GLU 299		PROT
	ATOM	631 C GLU 299	36.057 24.932 13.475 1.00 38.89	PROT
	ATOM	632 O GLU 299	37.129 24.342 13.316 1.00 48.67	PROT
	ATOM	633 N ASP 300	35.528 25.186 14.671 1.00 36.49	PROT
	ATOM	634 CA ASP 300	36.201 24.805 15.906 1.00 29.96	PROT
25	ATOM	635 CB ASP 300	35.455 25.391 17.111 1.00 5.33	PROT
	ATOM	636 CG ASP 300	35.830 26.853 17.378 1.00 19.10	PROT
	ATOM	637 OD1 ASP 300	36.491 27.473 16.518 1.00 27.28	PROT
	ATOM	638 OD2 ASP 300	35,470 27,396 18,444 1.00 23,55	PROT
	ATOM	639 C ASP 300	36.380 23.294 16.054 1.00 25.88	PROT
30	ATOM	640 O ASP 300	37.441 22.845 16.484 1.00 19.03	PROT
	ATOM	641 N GLN 301	35.360 22.516 15.689 1.00 6.29	PROT
	ATOM	642 CA GLN 301	35.432 21.055 15.769 1.00 9.51	PROT
	ATOM	643 CB GLN 301	34.170 20.421 15.183 1.00 18.27	PROT
	ATOM	644 CG GLN 301	32.886 20.813 15.875 1.00 28.72	PROT
35	ATOM	645 CD GLN 301	31.676 20.155 15.243 1.00 17.63	PROT
	ATOM	646 OE1 GLN 301	31.689 19.823 14.060 1.00 30.65	PROT
	ATOM	647 NE2 GLN 301	30.625 19.965 16.027 1.00 30.44	PROT
	ATOM	648 C GLN 301	36.646 20.491 15.020 1.00 15.48	PROT
	ATOM	649 O GLN 301	37.333 19.584 15.500 1.00 21.96	PROT
40	ATOM	650 N ILE 302	36.891 21.014 13.825 1.00 24.00	PROT
	ATOM	651 CA ILE 302	38.011 20.555 13.026 1.00 28.84	PROT
	ATOM	652 CB ILE 302	37.930 21.112 11.607 1.00 33.13	PROT
	ATOM	653 CG2 ILE 302	39.147 20.690 10.813 1.00 37.90	PROT
	ATOM	654 CG1 ILE 302	36.656 20.610 10.941 1.00 29.63	PROT
45	ATOM	655 CD1 ILE 302	36.296 21.356 9.698 1.00 32.99	PROT
	ATOM	656 C ILE 302	39.308 21.014 13.670 1.00 28.73	PROT
	ATOM	657 O ILE 302	40.219 20.219 13.895 1.00 36.02	PROT
	ATOM	658 N ILE 303	39.396 22.304 13.968 1.00 25.04	PROT
	ATOM	659 CA ILE 303	40.590 22.817 14.603 1.00 24.27	PROT
50	ATOM	660 CB ILE 303	40.414 24.270 15.054 1.00 20.89	PROT
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	ATOM	661 CG2 ILE 303	41.686 24.740 15.744 1.00 32.38	PROT
	ATOM	662 CG1 ILE 303	40.079 25.158 13.849 1.00 18.88	PROT
	ATOM	663 CD1 ILE 303	40.298 26.648 14.079 1.00 5.31	PROT
	ATOM		0.861 21.948 15.825 1.00 26.92	PROT
5	ATOM		1.963 21.440 15.997 1.00 31.32	PROT
	ATOM		39.843 21.763 16.659 1.00 11.00	PROT
	ATOM	667 CA LEU 304	39.983 20.953 17.854 1.00 7.21	PROT
	ATOM	668 CB LEU 304	38.663 20.886 18.613 1.00 2.00	PROT
	ATOM	669 CG LEU 304	38.633 21.511 20.012 1.00 8.04	PROT
10	ATOM	670 CD1 LEU 304	39.383 22.812 19.997 1.00 2.00	PROT
	ATOM	671 CD2 LEU 304	37.188 21.729 20.472 1.00 4.99	PROT
	ATOM		40.441 19.554 17.507 1.00 4.64	PROT
	ATOM		41.368 19.032 18.119 1.00 14.88	PROT
	ATOM		39.807 18.953 16.510 1.00 4.55	PROT
15	ATOM	675 CA LEU 305	40.140 17.590 16.093 1.00 7.03	PROT
	ATOM	676 CB LEU 305	39.099 17.098 15.104 1.00 3.70	PROT
	ATOM	677 CG LEU 305	38.164 16.054 15.691 1.00 10.31	PROT
	ATOM	678 CD1 LEU 305	36.744 16.340 15.245 1.00 2.00	PROT
0.0	ATOM	679 CD2 LEU 305	38.629 14.665 15.260 1.00 9.42	PROT
20	ATOM		41.527 17.418 15.483 1.00 10.17	PROT
	ATOM		42.174 16.374 15.651 1.00 7.58	PROT
	ATOM		41.975 18.442 14.765 1.00 9.98	PROT
	ATOM	683 CA LYS 306	43.283 18.408 14.127 1.00 9.14 43.409 19.558 13.131 1.00 18.85	PROT
25	ATOM	684 CB LYS 306		PROT
25	ATOM	685 CG LYS 306	42.815 19.270 11.763 1.00 25.44 42.198 20.529 11.178 1.00 29.07	PROT
	ATOM	686 CD LYS 306 687 CE LYS 306	42.698 20.808 9.774 1.00 37.81	PROT PROT
	ATOM	688 NZ LYS 306	43.867 19.964 9.403 1.00 30.48	PROT
	ATOM ATOM		44.376 18.522 15.175 1.00 7.31	PROT
30	ATOM		45.439 17.919 15.048 1.00 16.95	PROT
30	ATOM		44.097 19.295 16.218 1.00 12.67	PROT
	ATOM	692 CA GLY 307	45.062 19.484 17.279 1.00 7.25	PROT
	ATOM		45.297 18.269 18.150 1.00 15.08	PROT
	ATOM		46.441 17.972 18.488 1.00 20.11	PROT
35	ATOM		44.225 17.552 18.481 1.00 8.29	PROT
55	ATOM	696 CA CYS 308	44.286 16.380 19.364 1.00 3.44	PROT
	ATOM	697 CB CYS 308	43.097 16.402 20.326 1.00 14.26	PROT
	ATOM	698 SG CYS 308	41.539 15.750 19.634 1.00 21.83	PROT
	ATOM		44.344 14.995 18.738 1.00 8.37	PROT
40	ATOM		44.502 13.997 19.453 1.00 10.98	PROT
	ATOM		44.202 14.916 17.420 1.00 10.83	PROT
	ATOM	702 CA CYS 309	44.236 13.625 16.752 1.00 3.22	PROT
	ATOM	703 CB CYS 309	44.240 13.831 15.240 1.00 15.79	PROT
	ATOM	704 SG CYS 309	43.683 12.402 14.319 1.00 25.54	PROT
45	ATOM		45.439 12.767 17.193 1.00 2.00	PROT
	ATOM		45.251 11.722 17.807 1.00 12.28	PROT
	ATOM		46.663 13.205 16.900 1.00 2.00	<b>PROT</b>
	ATOM	708 CA MET 310	47.858 12.446 17.286 1.00 2.00	PROT
	ATOM	709 CB MET 310	49.122 13.171 16.860 1.00 2.00	PROT
50	ATOM	710 CG MET 310	49.975 12.422 15.880 1.00 5.92	PROT

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PROT
                               50.481 10.805 16.368 1.00 22.47
    ATOM
            711 SD MET 310
                               52.140 11.112 16.808 1.00 20.84
                                                             PROT
            712 CE MET 310
    ATOM
                              47.941 12.239 18.793 1.00 11.95
                                                             PROT
            713 C MET 310
    ATOM
                              48.455 11.220 19.270 1.00 15.53
                                                             PROT
            714 O MET 310
    ATOM
                              47.463 13.225 19.542 1.00 6.79
                                                            PROT
    ATOM
            715 N GLU 311
                               47.493 13.139 20.979 1.00 2.00
                                                             PROT
            716 CA GLU 311
    ATOM
                               46.932 14.427 21.581 1.00 6.42
                                                             PROT
            717 CB GLU 311
    ATOM
            718 CG GLU 311
                               47.880 15.619 21.436 1.00 8.40
                                                             PROT
    ATOM
                               47.236 16.940 21.820 1.00 14.10
                                                             PROT
    ATOM
            719 CD GLU 311
                               46.157 16.895 22.434 1.00 16.54
                                                             PROT
10
    ATOM
            720 OE1 GLU 311
                               47.795 18.020 21.515 1.00 4.09
                                                             PROT
    ATOM
            721 OE2 GLU 311
                              46.683 11.923 21.406 1.00 7.80
    ATOM
            722 C GLU 311
                                                            PROT
                              47.195 11.026 22.067 1.00 14.07
    ATOM
            723 O GLU 311
                                                            PROT
            724 N ILE 312
                             45.425 11.873 21.001 1.00 2.00
                                                           PROT
    ATOM
            725 CA ILE 312
                              44.574 10.752 21.371 1.00 3.60
                                                            PROT
15
    ATOM
            726 CB ILE 312
                              43.114 11.013 20.947 1.00 2.00
                                                            PROT
    ATOM
                               42.277 9.769 21.145 1.00 2.00
                                                            PROT
            727 CG2 ILE 312
    ATOM
                               42.579 12.221 21.727 1.00 2.00
                                                            PROT
    ATOM
            728 CG1 ILE 312
                               41.118 12.555 21.495 1.00 2.00
                                                            PROT
    ATOM
            729 CD1 ILE 312
                             45.049 9.437 20.760 1.00 8.32
                                                           PROT
            730 C ILE 312
20
    ATOM
            731 O ILE 312
                             44.918 8.373 21.370 1.00 5.58
                                                           PROT
    ATOM
                              45.615 9.501 19.563 1.00 3.98
                                                            PROT
           732 N MET 313
    ATOM
                               46.054 8.282 18.905 1.00 8.91
                                                            PROT
            733 CA MET 313
    ATOM
                               46.455 8.572 17.462 1.00 25.71
                                                             PROT
            734 CB MET 313
    ATOM
            735 CG MET 313
                                                             PROT
                               45.430 8.111 16.431 1.00 22.86
25
    ATOM
                               45.955 8.430 14.736 1.00 20.60
            736 SD MET 313
                                                             PROT
    ATOM
            737 CE MET 313
                               45.412 10.055 14.534 1.00 14.95
                                                             PROT
    ATOM
            738 C MET 313
                              47.211 7.634 19.635 1.00 12.95
                                                            PROT
    ATOM
                              47.213 6.426 19.857 1.00 22.09
                                                            PROT
            739 O MET 313
    ATOM
                              48.190 8.442 20.021 1.00 10.79
                                                            PROT
            740 N SER 314
    ATOM
30
                               49.354 7.935 20.719 1.00 2.00
                                                            PROT
            741 CA SER 314
    ATOM
                                                            PROT
                               50.399 9.042 20.816 1.00 7.24
    ATOM
            742 CB SER 314
                               50.453 9.815 19.619 1.00 10.89
                                                             PROT
    ATOM
            743 OG SER 314
            744 C SER 314
                              48.991 7.399 22.105 1.00 8.64
                                                           PROT
    ATOM
                                                           PROT
            745 O SER 314
                              49.559 6.392 22.558 1.00 5.72
35
    A·TOM
                              48.050 8.062 22.782 1.00 2.00
           746 N LEU 315
                                                            PROT
    ATOM
           747 CA LEU 315
                               47.628 7.605 24.104 1.00 2.00
                                                            PROT
    ATOM
                               46.521 8.502 24.671 1.00 2.95
                                                            PROT
            748 CB LEU 315
    ATOM
                               45.831 8.096 25.992 1.00 2.00
                                                            PROT
    ATOM
            749 CG LEU 315
                                46.876 7.845 27.072 1.00 2.54
                                                             PROT
            750 CD1 LEU 315
40
    ATOM
                                44,865 9.182 26.444 1.00 2.00
                                                             PROT
            751 CD2 LEU 315
    ATOM
                              47.107 6.182 23.945 1.00 3.25
                                                            PROT
    ATOM
            752 C LEU 315
                              47.568 5.253 24.603 1.00 2.00
            753 O LEU 315
                                                            PROT
    ATOM
                               46.157 6.010 23.039 1.00 7.28
                                                            PROT
    ATOM
            754 N ARG 316
                               45.588 4.691 22.808 1.00 13.31
                                                             PROT
45
    ATOM
            755 CA ARG 316
                               44.551 4.758 21.693 1.00 11.11
                                                             PROT
    ATOM
            756 CB ARG 316
            757 CG ARG 316
                               43.545 5.872 21.887 1.00 10.55
                                                             PROT
    ATOM
                               42.354 5.639 21.012 1.00 10.09
    ATOM
            758 CD ARG 316
                                                             PROT
    ATOM
            759 NE ARG 316
                               41.131 6.149 21.605 1.00 12.29
                                                             PROT
                               39.955 6.127 20.994 1.00 6.99
    ATOM
            760 CZ ARG 316
                                                             PROT
50
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	ATOM	761 NH1 ARG 316	38.880 6.608 21.595 1.00 19.32 39.853 5.619 19.778 1.00 17.16	PROT PROT
	ATOM	762 NH2 ARG 316	39.853 5.619 19.778 1.00 17.16 46.666 3.686 22.458 1.00 10.10	PROT
	ATOM	763 C ARG 316 764 O ARG 316	46.549 2.508 22.753 1.00 10.10	PROT
5	ATOM ATOM		47.723 4.148 21.819 1.00 6.51	PROT
3	_	765 N ALA 317 766 CA ALA 317	48.801 3.243 21.474 1.00 11.04	PROT
	ATOM	767 CB ALA 317	49.749 3.902 20.487 1.00 16.13	PROT
	ATOM ATOM	767 CB ALA 317	49.539 2.910 22.753 1.00 12.70	PROT
	ATOM	769 O ALA 317	49.822 1.755 23.033 1.00 23.09	PROT
10	ATOM	770 N ALA 318	49.832 3.943 23.534 1.00 14.79	PROT
10	ATOM	771 CA ALA 318	50.567 3.779 24.776 1.00 8.38	PROT
	ATOM	771 CA ALA 318	50.727 5.122 25.448 1.00 11.75	PROT
	ATOM	772 CB ALA 318	49.941 2.786 25.741 1.00 10.30	PROT
	ATOM	774 O ALA 318	50.585 1.824 26.165 1.00 8.48	PROT
15	ATOM	774 O ALA 318 775 N VAL 319	48.680 3.011 26.083 1.00 7.87	PROT
13	ATOM	776 CA VAL 319	48.002 2.131 27.027 1.00 9.64	PROT
	ATOM	777 CB VAL 319	46.579 2.622 27.334 1.00 2.57	PROT
	ATOM	778 CG1 VAL 319	46.644 3.929 28.127 1.00 5.09	PROT
	ATOM	779 CG2 VAL 319	45.807 2.823 26.043 1.00 5.15	PROT
20	ATOM	780 C VAL 319	47.930 0.695 26.541 1.00 11.68	PROT
	ATOM	781 O VAL 319	47,440 -0.171 27.254 1.00 16.32	PROT
	ATOM	782 N ARG 320	48.415 0.444 25.329 1.00 16.40	PROT
	ATOM	783 CA ARG 320	48,405 -0.902 24.767 1.00 13.20	PROT
	ATOM	784 CB ARG 320	47.736 -0.918 23.393 1.00 2.00	PROT
25	ATOM	785 CG ARG 320	46.310 -0.405 23.420 1.00 14.07	PROT
	<b>ATOM</b>	786 CD ARG 320	45.283 -1.460 23.035 1.00 19.69	PROT
	ATOM	787 NE ARG 320	44.168 -0.868 22.292 1.00 36.52	PROT
	ATOM	788 CZ ARG 320	42.912 -1.313 22.322 1.00 47.43	PROT
	ATOM	789 NH1 ARG 320	41.966 -0.705 21.609 1.00 43.57	PROT
30	<b>ATOM</b>	790 NH2 ARG 320	42.596 -2.367 23.061 1.00 49.93	PROT
	ATOM	791 C ARG 320	49.835 -1.391 24.662 1.00 15.45	PROT
	ATOM	792 O ARG 320	50.167 -2.218 23.809 1.00 24.78	PROT
	ATOM	793 N TYR 321	50.684 -0.860 25.537 1.00 13.68	PROT
	ATOM	794 CA TYR 321	52.085 -1.258 25.572 1.00 18.80	PROT
35	ATOM	795 CB TYR 321	52.925 -0.208 26.295 1.00 9.64	PROT
	ATOM	796 CG TYR 321	54.313 -0.685 26.622 1.00 11.20	PROT
	ATOM	797 CD1 TYR 321	55.211 -1.005 25.612 1.00 2.00	PROT
	ATOM	798 CE1 TYR 321	56.483 -1.461 25.906 1.00 9.63	PROT
	ATOM	799 CD2 TYR 321	54.727 -0.834 27.943 1.00 18.93	PROT
40	ATOM	800 CE2 TYR 321	56.003 -1.293 28.250 1.00 19.49	PROT
	ATOM	801 CZ TYR 321	56.874 -1.604 27.225 1.00 14.75	PROT
	ATOM	802 OH TYR 321	58.137 -2.053 27.518 1.00 22.96	PROT
	ATOM	803 C TYR 321	52.209 -2.607 26.287 1.00 19.74	PROT
15	ATOM	804 O TYR 321	51.483 -2.889 27.242 1.00 31.56	PROT
45	ATOM	805 N ASP 322	53.136 -3.435 25.823 1.00 26.35	PROT
	ATOM	806 CA ASP 322	53.346 -4.759 26.392 1.00 22.38	PROT
	ATOM	807 CB ASP 322 808 CG ASP 322	52.982 -5.814 25.353 1.00 33.63 52.601 -7.128 25.970 1.00 40.70	PROT PROT
	ATOM ATOM	808 CG ASP 322 809 OD1 ASP 322	51.539 -7.658 25.591 1.00 40.70	PROT
50	ATOM	810 OD2 ASP 322	53.358 -7.628 26.826 1.00 38.91	PROT
JU	ATOM	010 ODZ ASP 322	-1.020 20.020 1.00 30.71	1101

	ATOM	811 C ASP 322	54.800 -4.928 26.776 1.00 23.51	PROT
	ATOM	812 O ASP 322	55.683 -4.844 25.924 1.00 37.80	PROT
	ATOM	813 N PRO 323	55.076 -5.160 28.066 1.00 24.06	PROT
	ATOM	814 CD PRO 323	54.130 -5.258 29.187 1.00 19.35	PROT
5	ATOM	815 CA PRO 323	56.462 -5.339 28.507 1.00 23.60	PROT
	ATOM	816 CB PRO 323	56.390 -5.121 30.007 1.00 3.90	PROT
	ATOM	817 CG PRO 323	55.031 -5.570 30.360 1.00 14.06	PROT
	ATOM	818 C PRO 323	56.949 -6.736 28.151 1.00 21.79	PROT
	ATOM	819 O PRO 323	58.149 -7.003 28.119 1.00 27.28	PROT
10	ATOM	820 N GLU 324	56.009 -7.633 27.889 1.00 37.63	PROT
	ATOM	821 CA GLU 324	56.366 -8.993 27.524 1.00 42.63	PROT
	ATOM	822 CB GLU 324	55.133 -9.885 27.551 1.00 37.58	PROT
	ATOM	823 C GLU 324	56.971 -8.956 26.124 1.00 43.28	PROT
	ATOM	824 O GLU 324	58.154 -9.239 25.938 1.00 43.14	PROT
15	ATOM	825 N SER 325	56.153 -8.586 25.142 1.00 31.72	PROT
	ATOM	826 CA SER 325	56.607 -8.508 23.765 1.00 30.34	PROT
	ATOM	827 CB SER 325	55.413 -8.522 22.814 1.00 17.63	PROT
	ATOM	828 OG SER 325	54.356 -7.729 23.315 1.00 31.90	PROT
	ATOM	829 C SER 325	57.441 -7.257 23.519 1.00 31.94	PROT
20	ATOM	830 O SER 325	58.146 -7.169 22.513 1.00 45.47	PROT
	ATOM	831 N GLU 326	57.359 -6.289 24.429 1.00 31.10	PROT
	ATOM	832 CA GLU 326	58.119 -5.050 24.281 1.00 31.43	PROT
	ATOM	833 CB GLU 326	59.598 -5.382 24.091 1.00 30.39	PROT
	ATOM	834 CG GLU 326	60.552 -4.342 24.612 1.00 35.00	PROT
25	ATOM	835 CD GLU 326	61.738 -4.965 25.304 1.00 29.12	PROT
	ATOM	836 OE1 GLU 326	61.525 -5.579 26.370 1.00 39.21	PROT
	ATOM	837 OE2 GLU 326	62.872 -4.844 24.788 1.00 29.11	PROT
	ATOM	838 C GLU 326	57.605 -4.283 23.063 1.00 28.37	PROT
	ATOM	839 O GLU 326	58.382 -3.677 22.321 1.00 26.51	PROT
30	ATOM	840 N THR 327	56.290 -4.301 22.873 1.00 23.71	PROT
	ATOM	841 CA THR 327	55.674 -3.648 21.720 1.00 22.11	PROT
	ATOM	842 CB THR 327	55.298 -4.705 20.652 1.00 28.08	PROT
	ATOM	843 OG1 THR 327	54.226 -5.524 21.145 1.00 16.87	PROT
	ATOM	844 CG2 THR 327	56,494 -5.597 20.340 1.00 24.03	PROT
35	ATOM	845 C THR 327	54,420 -2.824 22.046 1.00 22.42	PROT
	ATOM	846 O THR 327	53.928 -2.830 23.172 1.00 17.50	PROT
	ATOM	847 N LEU 328	53.914 -2.122 21.038 1.00 17.28	PROT
	ATOM	848 CA LEU 328	52.728 -1.285 21.171 1.00 14.83	PROT
	ATOM	849 CB LEU 328	53.065 0.157 20.806 1.00 15.27	PROT
40	ATOM	850 CG LEU 328	53.693 1.036 21.879 1.00 10.50	PROT
. •	ATOM	851 CD1 LEU 328	54.137 2.336 21.254 1.00 16.75	PROT
	ATOM	852 CD2 LEU 328	52.682 1.285 22.979 1.00 20.19	PROT
	ATOM	853 C LEU 328	51.687 -1.804 20.198 1.00 18.16	PROT
	ATOM	854 O LEU 328	52.035 -2.508 19.254 1.00 23.88	PROT
45	ATOM	855 N THR 329	50.421 -1.450 20.402 1.00 9.40	PROT
	ATOM	856 CA THR 329	49.389 -1.920 19.495 1.00 8.26	PROT
	ATOM	857 CB THR 329	48.460 -2.888 20.199 1.00 8.67	PROT
	ATOM	858 OG1 THR 329	49.213 -4.052 20.577 1.00 13.23	PROT
	ATOM	859 CG2 THR 329	47.308 -3.289 19.270 1.00 2.00	PROT
50	ATOM	860 C THR 329	48.569 -0.841 18.800 1.00 16.65	PROT
		555 C 1111C 527	.5.557 5.5	- 450 1

	ATOM	861 O THR 329	47.726 -0.158 19.406 1.00 17.20	PROT
	ATOM	862 N LEU 330	48.808 -0.725 17.495 1.00 21.56	PROT
	ATOM	863 CA LEU 330	48.138 0.258 16.655 1.00 20.95	PROT
	ATOM	864 CB LEU 330	49.106 0.676 15.539 1.00 17.36	PROT
5	ATOM	865 CG LEU 330	50.570 0.797 16.028 1.00 12.86	PROT
	ATOM	866 CD1 LEU 330	51.531 0.521 14.898 1.00 10.10	PROT
	ATOM	867 CD2 LEU 330	50.830 2.180 16.600 1.00 2.00	PROT
	<b>ATOM</b>	868 C LEU 330	46.803 -0.258 16.097 1.00 21.35	PROT
	<b>ATOM</b>	869 O LEU 330	46.655 -1.444 15.791 1.00 21.93	PROT
10	ATOM	870 N ASN 331	45.834 0.648 15.987 1.00 27.76	PROT
	ATOM	871 CA ASN 331	44.487 0.338 15.498 1.00 28.09	PROT
	ATOM	872 CB ASN 331	44.460 0.275 13.971 1.00 24.95	PROT
	ATOM	873 CG ASN 331	43.074 0.540 13.397 1.00 33.45	PROT
	ATOM	874 OD1 ASN 331	42.512 -0.305 12.701 1.00 38.21	PROT
15	ATOM	875 ND2 ASN 331	42.522 1.715 13.680 1.00 24.73	PROT
	ATOM	876 C ASN 331	43.946 -0.967 16.075 1.00 32.03	PROT
	ATOM	877 O ASN 331	43.166 -1.668 15.431 1.00 35.49	PROT
	ATOM	878 N GLY 332	44.357 -1.282 17.299 1.00 40.24	PROT
	ATOM	879 CA GLY 332	43.894 -2.495 17.941 1.00 38.04	PROT
20	ATOM	880 C GLY 332	44.009 -3.665 16.998 1.00 40.09	PROT
	ATOM	881 O GLY 332	43.001 -4.225 16.563 1.00 45.79	PROT
	ATOM	882 N GLU 333	45.249 -4.013 16.664 1.00 41.60	PROT
	ATOM	883 CA GLU 333	45.539 -5.126 15.763 1.00 36.28	PROT
	ATOM	884 CB GLU 333	44.752 -4.978 14.454 1.00 46.39	PROT
25	ATOM	885 CG GLU 333	44.745 -3.580 13.862 1.00 58.03	PROT
	ATOM	886 CD GLU 333	43.883 -3.485 12.610 1.00 67.00	PROT
	ATOM	887 OE1 GLU 333	44.446 -3.282 11.511 1.00 67.51	PROT
	ATOM	888 OE2 GLU 333	42.644 -3.615 12.727 1.00 71.01	PROT
	ATOM	889 C GLU 333	47.027 -5.266 15.446 1.00 33.13	PROT
30	ATOM	890 O GLU 333	47.563 -6.366 15.486 1.00 27.97	PROT
	ATOM	891 N MET 334	47.692 -4.152 15.143 1.00 27.00	PROT
	ATOM	892 CA MET 334	49.111 -4.188 14.798 1.00 29.83	PROT
	ATOM	893 CB MET 334	49.416 -3.159 13.699 1.00 26.04	PROT
	ATOM	894 CG MET 334	50.561 -3.588 12.765 1.00 28.06	PROT
35	ATOM	895 SD MET 334	51.263 -2.273 11.736 1.00 28.46	PROT
-	ATOM	896 CE MET 334	50.021 -2.123 10.497 1.00 22.48	PROT
	ATOM	897 C MET 334	50.087 -3.995 15.959 1.00 33.52	PROT
	ATOM	898 O MET 334	50.071 -2.962 16.631 1.00 35.81	PROT
	ATOM	899 N ALA 335	50.942 -4.996 16.171 1.00 27.46	PROT
40	ATOM	900 CA ALA 335	51.948 -4.976 17.234 1.00 29.69	PROT
	ATOM	901 CB ALA 335	51.966 -6.314 17.965 1.00 12.67	PROT
	ATOM	902 C ALA 335	53.336 -4.682 16.662 1.00 31.74	PROT
	ATOM	903 O ALA 335	53.943 -5.530 16.009 1.00 43.66	PROT
	ATOM	904 N VAL 336	53.848 -3.489 16.923 1.00 23.98	PROT
45	ATOM	905 CA VAL 336	55.151 -3.118 16.405 1.00 21.32	PROT
73	ATOM	905 CA VAL 336	55.028 -1.873 15.504 1.00 17.37	PROT
	ATOM	907 CG1 VAL 336	53.945 -2.104 14.462 1.00 14.88	PROT
	ATOM	907 CG1 VAL 336 908 CG2 VAL 336	54.686 -0.648 16.339 1.00 15.53	PROT
	ATOM	909 C VAL 336	56.150 -2.852 17.526 1.00 22.72	PROT
50	ATOM	910 O VAL 336	55.763 -2.540 18.651 1.00 25.15	PROT
20	A I OIVI	210 O AWP 330	JJ./UJ -Z.J4U 10.UJ1 1.UU 2J.1J	IVOI

	ATOM	911 N THR 337	57.435 -3.001 17.220 1.00 19.21	PROT
	ATOM	912 CA THR 337	58.476 -2.765 18.205 1.00 20.31	PROT
	ATOM	913 CB THR 337	59.752 -3.578 17.884 1.00 14.76	PROT
	ATOM	914 OG1 THR 337	59.957 -3.616 16.467 1.00 16.43	PROT
5	ATOM	915 CG2 THR 337	59.615 -4.995 18.393 1.00 7.08	PROT
	ATOM	916 C THR 337	58.785 -1.272 18.157 1.00 24.20	PROT
	ATOM	917 O THR 337	58.322 -0.591 17.245 1.00 28.05	PROT
	ATOM	918 N ARG 338	59.548 -0.766 19.134 1.00 27.55	PROT
	ATOM	919 CA ARG 338	59.917 0.655 19.197 1.00 16.80	PROT
10	ATOM	920 CB ARG 338	60.757 0.942 20.446 1.00 17.04	PROT
	ATOM	921 CG ARG 338	61.687 2.149 20.303 1.00 9.79	PROT
	ATOM	922 CD ARG 338	62.666 2.276 21.458 1.00 2.00	PROT
	ATOM	923 NE ARG 338	61.994 2.128 22.739 1.00 20.70	PROT
	ATOM	924 CZ ARG 338	61.897 3.083 23.657 1.00 12.04	PROT
15	ATOM	925 NH1 ARG 338	61.261 2.840 24.784 1.00 27.11	PROT
	ATOM	926 NH2 ARG 338	62.436 4.272 23.459 1.00 22.23	PROT
	ATOM	927 C ARG 338	60.702 1.085 17.968 1.00 21.26	PROT
	ATOM	928 O ARG 338	60.338 2.049 17.295 1.00 16.40	PROT
	ATOM	929 N GLY 339	61.792 0.374 17.693 1.00 31.57	PROT
20	ATOM	930 CA GLY 339	62.609 0.696 16.540 1.00 32.42	PROT
	ATOM	931 C GLY 339	61.816 0.534 15.254 1.00 30.08	PROT
	ATOM	932 O GLY 339	61.932 1.342 14.328 1.00 25.82	PROT
	ATOM	933 N GLN 340	61.008 -0.520 15.192 1.00 16.60	PROT
	ATOM	934 CA GLN 340	60.191 -0.768 14.012 1.00 14.08	PROT
25	ATOM	935 CB GLN 340	59.199 -1.884 14.301 1.00 5.73	PROT
	ATOM	936 CG GLN 340	58.849 -2.697 13.100 1.00 16.15	PROT
	ATOM	937 CD GLN 340	58.577 -4.141 13.442 1.00 22.46	PROT
	ATOM	938 OE1 GLN 340	57.767 -4.450 14.316 1.00 30.45	PROT
••	ATOM	939 NE2 GLN 340	59.254 -5.040 12.749 1.00 34.19	PROT PROT
30	ATOM	940 C GLN 340	59.452 0.521 13.632 1.00 22.07	PROT
	ATOM	941 O GLN 340	59.707 1.103 12.576 1.00 21.13	PROT
	ATOM	942 N LEU 341	58.561 0.976 14.518 1.00 27.88	
	ATOM	943 CA LEU 341	57.778 2.197 14.306 1.00 21.82	PROT PROT
2.5	ATOM	944 CB LEU 341	56.813 2.418 15.483 1.00 10.20 55.930 3.682 15.534 1.00 16.27	PROT
35	ATOM	945 CG LEU 341	55.930 3.682 15.534 1.00 16.27 54.777 3.618 14.518 1.00 13.27	PROT
	ATOM	946 CD1 LEU 341		PROT
	ATOM	947 CD2 LEU 341		PROT
	ATOM	948 C LEU 341	58.683 3.413 14.138 1.00 13.98 58.315 4.386 13.486 1.00 7.94	PROT
40	ATOM	949 O LEU 341		PROT
40	ATOM	950 N LYS 342	59.867 3.361 14.734 1.00 11.48 60.804 4.465 14.613 1.00 17.77	PROT
	ATOM	951 CA LYS 342 952 CB LYS 342	62.063 4.213 15.459 1.00 13.58	PROT
	ATOM		63.219 5.173 15.140 1.00 13.27	PROT
	ATOM	953 CG LYS 342 954 CD LYS 342	64.173 5.358 16.319 1.00 5.44	PROT
15	ATOM			PROT
45	ATOM	955 CE LYS 342	64.500 6.829 16.546 1.00 5.47 65.721 7.019 17.388 1.00 4.98	PROT
	ATOM	956 NZ LYS 342	61.184 4.579 13.141 1.00 19.97	PROT
	ATOM	957 C LYS 342 958 O LYS 342	60.939 5.595 12.501 1.00 20.34	PROT
	ATOM ATOM	958 O LYS 342 959 N ASN 343	61.764 3.510 12.605 1.00 26.88	PROT
50		960 CA ASN 343	62.196 3.470 11.219 1.00 22.34	PROT
50	ATOM	900 CA ASIN 343	02.170 3.470 11.219 1.00 22.34	1101

	ATOM	961 CB ASN 343	62.829 2.123 10.929 1.00 4.80	PROT
	ATOM	962 CG ASN 343	64.060 1.894 11.758 1.00 18.77	PROT
	ATOM	963 ODI ASN 343	64.755 2.848 12.117 1.00 14.12	PROT
_	ATOM	964 ND2 ASN 343	64.340 0.634 12.083 1.00 12.72	PROT PROT
5	ATOM	965 C ASN 343	61.091 3.736 10.224 1.00 20.40 61.309 4.417 9.232 1.00 20.76	PROT
	ATOM ATOM	966 O ASN 343 967 N GLY 344	61.309 4.417 9.232 1.00 20.76 59.908 3.200 10.494 1.00 12.62	PROT
	ATOM	967 N GLY 344 968 CA GLY 344	58.775 3.382 9.603 1.00 6.27	PROT
	ATOM	969 C GLY 344	58.229 4.796 9.451 1.00 14.56	PROT
10	ATOM	970 O GLY 344	57.177 4.972 8.826 1.00 13.30	PROT
10	ATOM	971 N GLY 345	58.902 5.795 10.030 1.00 16.51	PROT
	ATOM	971 N GL1 343 972 CA GLY 345	58.439 7.166 9.869 1.00 20.04	PROT
	ATOM	972 CA GL1 343 973 C GLY 345	58.248 8.112 11.046 1.00 25.64	PROT
	ATOM	974 O GLY 345	58.243 9.331 10.849 1.00 23.32	PROT
15	ATOM	974 O GET 343 975 N LEU 346	58.099 7.588 12.260 1.00 22.22	PROT
15	ATOM	976 CA LEU 346	57.874 8.449 13.415 1.00 14.94	PROT
	ATOM	977 CB LEU 346	57.070 7.700 14.474 1.00 3.92	PROT
	ATOM	978 CG LEU 346	55.566 7.538 14.193 1.00 5.92	PROT
	ATOM	979 CD1 LEU 346	54.938 6.796 15.355 1.00 2.00	PROT
20	ATOM	980 CD2 LEU 346	54.884 8.885 13.973 1.00 2.00	PROT
	ATOM	981 C LEU 346	59.126 9.042 14.041 1.00 14.60	PROT
	ATOM	982 O LEU 346	59.102 10.153 14.554 1.00 17.36	PROT
	ATOM	983 N GLY 347	60.226 8.312 14.001 1.00 12.09	PROT
	ATOM	984 CA GLY 347	61.455 8.828 14.581 1.00 15.62	PROT
25	ATOM	985 C GLY 347	61.439 8.963 16.090 1.00 6.31	PROT
	ATOM	986 O GLY 347	60.865 8.141 16.790 1.00 13.15	PROT
	ATOM	987 N VAL 348	62.076 10.011 16.592 1.00 13.74	PROT
	ATOM	988 CA VAL 348	62.141 10.259 18.030 1.00 10.13	PROT
	ATOM	989 CB VAL 348	62.757 11.646 18.342 1.00 9.26	PROT
30	ATOM	990 CG1 VAL 348	61.867 12.752 17.794 1.00 2.00	PROT
	ATOM	991 CG2 VAL 348	62.942 11.802 19.836 1.00 2.00	PROT
	ATOM	992 C VAL 348	60.763 10.216 18.650 1.00 6.61	PROT
	ATOM	993 O VAL 348	60.619 10.066 19.862 1.00 3.12	PROT
	ATOM	994 N VAL 349	59.746 10.358 17.816 1.00 5.51	PROT
35	ATOM	995 CA VAL 349	58.386 10.342 18.306 1.00 2.00	PROT
	ATOM	996 CB VAL 349	57.421 10.886 17.260 1.00 4.46	PROT
	ATOM	997 CG1 VAL 349	56.001 10.578 17.656 1.00 2.00	PROT
	ATOM	998 CG2 VAL 349	57.623 12.387 17.122 1.00 2.00	PROT
40	ATOM	999 C VAL 349	57.995 8.933 18.687 1.00 9.15	PROT
40	ATOM	1000 O VAL 349	57.284 8.726 19.664 1.00 15.02	PROT
	ATOM	1001 N SER 350	58.446 7.943 17.933 1.00 7.42	PROT PROT
	ATOM	1002 CA SER 350 1003 CB SER 350	58.087 6.590 18.315 1.00 12.87 58.695 5.561 17.382 1.00 9.48	PROT
	ATOM ATOM	1003 CB SER 350 1004 OG SER 350	58.529 4.269 17.931 1.00 10.82	PROT
15			58.529 4.269 17.931 1.00 10.82 58.628 6.364 19.717 1.00 15.55	PROT
45	ATOM ATOM	1005 C SER 350 1006 O SER 350	57.963 5.761 20.558 1.00 25.88	PROT
	ATOM	1006 O SER 350 1007 N ASP 351	59.838 6.863 19.950 1.00 16.38	PROT
	ATOM	1007 N ASP 331 1008 CA ASP 351	60.522 6.743 21.230 1.00 9.58	PROT
	ATOM	1008 CA ASI 351 1009 CB ASP 351	61.861 7.469 21.176 1.00 7.32	PROT
50	ATOM	1010 CG ASP 351	62.989 6.576 20.742 1.00 24.16	PROT
23	1110111	1010 00 1101 001	52.707 5.5.0 20.7.2 1.00 2 1.10	1

	ATOM		1.011 7.110 20.275 1.00 30.24	PROT
	ATOM		2.866 5.343 20.869 1.00 33.85	PROT
	ATOM	1013 C ASP 351 59.0		PROT
_	ATOM	1014 O ASP 351 59.6		PROT
5	ATOM		100 8.508 22.032 1.00 13.51	PROT
	ATOM		3.294 9.224 23.004 1.00 5.19	PROT
	ATOM		7.914 10.593 22.452 1.00 2.00	PROT
	ATOM		055 8.432 23.374 1.00 2.00 701 8.360 24.535 1.00 7.20	PROT PROT
10	ATOM			PROT
10	ATOM		201 7.049 22.677 1.00 5.90	PROT
	ATOM		468 6.626 21.381 1.00 5.87	PROT
	ATOM		113 6.049 21.732 1.00 2.00	PROT
	ATOM		349 7.831 20.428 1.00 3.91	PROT
15	ATOM ATOM		330 7.664 19.294 1.00 2.00	PROT
15	ATOM	1025 CDI ILE 353 55.5		PROT
	ATOM	1020 C ILE 353 54.8		PROT
	ATOM		544 5.131 23.122 1.00 19.57	PROT
	ATOM		.034 3.944 23.862 1.00 14.42	PROT
20	ATOM		256 3.270 23.209 1.00 3.70	PROT
20	ATOM		.890 2.141 22.284 1.00 9.42	PROT
	ATOM		7.427 2.401 20.995 1.00 12.33	PROT
	ATOM		7.912 0.822 22.727 1.00 15.63	PROT
	ATOM		.982 1.366 20.165 1.00 6.67	PROT
25	ATOM		.468 -0.224 21.900 1.00 16.53	PROT
	ATOM	1036 CZ PHE 354 57	002 0.053 20.620 1.00 11.61	PROT
	ATOM	1037 C PHE 354 57.3	322 4.346 25.307 1.00 18.55	PROT
	ATOM	1038 O PHE 354 56.	796 3.740 26.233 1.00 16.67	PROT
	ATOM	1039 N ASP 355 58.3		PROT
30	ATOM		.486 5.881 26.818 1.00 5.31	PROT
	ATOM		351 7.132 26.697 1.00 9.38	PROT
	ATOM		805 6.814 26.428 1.00 5.96	PROT
	ATOM		.112 5.683 26.016 1.00 8.53	PROT
	ATOM		.650 7.706 26.628 1.00 15.51	PROT
35	ATOM		252 6.199 27.659 1.00 10.27	PROT
	ATOM	1046 O ASP 355 57.2		PROT
	ATOM		224 6.726 27.014 1.00 4.18 .988 7.061 27.697 1.00 2.07	PROT
	ATOM ATOM		.988 7.061 27.697 1.00 2.07 .086 7.865 26.771 1.00 2.24	PROT PROT
40	ATOM		.694 8.229 27.266 1.00 3.11	PROT
40	ATOM		2.771 9.317 28.323 1.00 2.00	PROT
	ATOM		.877 8.709 26.086 1.00 2.00	PROT
	ATOM		281 5.786 28.091 1.00 9.17	PROT
	ATOM		331 5.644 29.221 1.00 14.77	PROT
45	ATOM		183 4.856 27.147 1.00 13.10	PROT
1.5	ATOM		.515 3.597 27.413 1.00 6.91	PROT
	ATOM		113 2.879 28.598 1.00 8.33	PROT
	ATOM		400 2.426 29.492 1.00 9.09	PROT
	ATOM		435 2.768 28.607 1.00 12.61	PROT
50	ATOM		5.112 2:091 29.692 1.00 10.53	PROT

	ATOM	1061 CB MET 358	57.626 2.153 29.498 1.00 5.45	PROT
	ATOM	1062 CG MET 358	58.138 1.507 28.210 1.00 15.15	PROT
	ATOM	1063 SD MET 358	59.971 1.352 28.113 1.00 17.63	PROT
	ATOM	1064 CE MET 358	60.445 3.023 27.774 1.00 20.56	PROT
5	ATOM	1065 C MET 358	55.714 2.809 30.972 1.00 15.08	PROT
	ATOM	1066 O MET 358	55.241 2.191 31.920 1.00 27.69	PROT
	ATOM	1067 N SER 359	55.875 4.125 30.984 1.00 20.67	PROT
	ATOM	1068 CA SER 359	55.551 4.924 32.158 1.00 19.72	PROT
	ATOM	1069 CB SER 359	55.831 6.398 31.861 1.00 19.98	PROT
10	ATOM	1070 OG SER 359	54.753 7.220 32.262 1.00 33.66	PROT
	ATOM	1071 C SER 359	54.115 4.757 32.656 1.00 22.67	PROT
	<b>ATOM</b>	1072 O SER 359	53.849 4.837 33.860 1.00 22.94	PROT
	<b>ATOM</b>	1073 N LEU 360	53.197 4.514 31.727 1.00 20.55	PROT
	ATOM	1074 CA LEU 360	51.785 4.360 32.054 1.00 17.01	PROT
15	ATOM	1075 CB LEU 360	50.934 4.578 30.802 1.00 2.60	PROT
	ATOM	1076 CG LEU 360	50.674 5.988 30.291 1.00 6.99	PROT
	<b>ATOM</b>	1077 CD1 LEU 360	49.589 5.935 29.236 1.00 4.15	PROT
	ATOM	1078 CD2 LEU 360	50.247 6.892 31.432 1.00 18.93	PROT
	<b>ATOM</b>	1079 C LEU 360	51.437 3.001 32.638 1.00 19.29	PROT
20	ATOM	1080 O LEU 360	50.319 2.802 33.102 1.00 27.53	PROT
	ATOM	1081 N SER 361	52.375 2.061 32.596 1.00 21.73	PROT
	ATOM	1082 CA SER 361	52.139 0.712 33.114 1.00 23.03	PROT
	ATOM	1083 CB SER 361	53.415 -0.130 33.027 1.00 25.89	PROT
	ATOM	1084 OG SER 361	53.645 -0.613 31.717 1.00 27.77	PROT
25	ATOM	1085 C SER 361	51.681 0.730 34.563 1.00 23.26	PROT
	ATOM	1086 O SER 361	50.720 0.046 34.929 1.00 18.73	PROT
	ATOM	1087 N SER 362	52.388 1.524 35.367 1.00 29.84	PROT
	ATOM	1088 CA SER 362	52.141 1.668 36.799 1.00 24.49	PROT
	<b>ATOM</b>	1089 CB SER 362	53.435 2.089 37.491 1.00 26.14	PROT
30	<b>ATOM</b>	1090 OG SER 362	53.917 3.305 36.949 1.00 25.03	PROT
	ATOM	1091 C SER 362	51.031 2.635 37.210 1.00 26.86	PROT
	<b>ATOM</b>	1092 O SER 362	50.797 2.831 38.404 1.00 39.63	PROT
	ATOM	1093 N PHE 363	50.361 3.251 36.240 1.00 20.94	PROT
	<b>ATOM</b>	1094 CA PHE 363	49.272 4.185 36.545 1.00 18.33	PROT
35	<b>ATOM</b>	1095 CB PHE 363	49.191 5.294 35.486 1.00 17.03	PROT
	ATOM	1096 CG PHE 363	50.171 6.407 35.706 1.00 22.73	PROT
	<b>ATOM</b>	1097 CD1 PHE 363	49.733 7.689 35.990 1.00 9.72	PROT
	ATOM	1098 CD2 PHE 363	51.545 6.167 35.659 1.00 24.77	PROT
	ATOM	1099 CE1 PHE 363	50.645 8.712 36.225 1.00 16.85	PROT
40	<b>ATOM</b>	1100 CE2 PHE 363	52.463 7.198 35.897 1.00 14.26	PROT
	ATOM	1101 CZ PHE 363	52.011 8.462 36.179 1.00 2.26	PROT
	<b>ATOM</b>	1102 C PHE 363	47.958 3.417 36.598 1.00 16.57	PROT
	ATOM	1103 O PHE 363	46.971 3.882 37.165 1.00 13.08	PROT
	ATOM	1104 N ASN 364	47.976 2.231 36.002 1.00 17.31	PROT
45	ATOM	1105 CA ASN 364	46.819 1.349 35.949 1.00 26.11	PROT
_	ATOM	1106 CB ASN 364	46.673 0.608 37.276 1.00 16.96	PROT
	ATOM	1107 CG ASN 364	47.402 -0.715 37.267 1.00 31.34	PROT
	ATOM	1108 OD1 ASN 364	46.965 -1.657 36.613 1.00 36.66	PROT
	ATOM	1109 ND2 ASN 364	48.527 -0.794 37.985 1.00 31.61	PROT
50	ATOM	1110 C ASN 364	45.527 2.060 35.594 1.00 18.22	PROT

	ATOM	1111 O ASN 364	44.522 1.923 36.286 1.00 23.17	PROT
	ATOM	1112 N LEU 365	45.567 2.803 34.491 1.00 13.10	PROT
	ATOM	1113 CA LEU 365	44.417 3.562 34.013 1.00 15.41	PROT
	ATOM	1114 CB LEU 365	44.833 4.483 32.861 1.00 16.55	PROT
5	ATOM	1115 CG LEU 365	45.762 5.653 33.181 1.00 19.56	PROT
	ATOM	1116 CD1 LEU 365	46.146 6.373 31.897 1.00 6.69	PROT
	ATOM	1117 CD2 LEU 365	45.067 6.602 34.128 1.00 15.69	PROT
	ATOM	1118 C LEU 365	43.328 2.624 33.520 1.00 12.07	PROT
	ATOM	1119 O LEU 365	43.620 1.534 33.043 1.00 19.81	PROT
10	<b>ATOM</b>	1120 N ASP 366	42.077 3.047 33.653 1.00 10.86	PROT
	<b>ATOM</b>	1121 CA ASP 366	40.942 2.263 33.180 1.00 8.96	PROT
	<b>ATOM</b>	1122 CB ASP 366	39.933 2.021 34.326 1.00 9.59	PROT
	<b>ATOM</b>	1123 CG ASP 366	39.300 3.306 34.859 1.00 21.78	PROT
	<b>ATOM</b>	1124 OD1 ASP 366	39.871 4.397 34.676 1.00 25.60	PROT
15	ATOM	1125 OD2 ASP 366	38.217 3.222 35.474 1.00 19.16	PROT
	ATOM	1126 C ASP 366	40.288 3.005 32.002 1.00 8.82	PROT
	<b>ATOM</b>	1127 O ASP 366	40.666 4.132 31.681 1.00 17.66	PROT
	<b>ATOM</b>	1128 N ASP 367	39.321 2.379 31.346 1.00 9.45	PROT
	ATOM	1129 CA ASP 367	38.668 3.023 30.218 1.00 11.11	PROT
20	ATOM	1130 CB ASP 367	37.457 2.205 29.769 1.00 20.67	PROT
	ATOM	1131 CG ASP 367	37.832 0.812 29.301 1.00 25.02	PROT
	<b>ATOM</b>	1132 OD1 ASP 367	39.040 0.525 29.158 1.00 21.06	PROT
	<b>ATOM</b>	1133 OD2 ASP 367	36.909 0.002 29.076 1.00 31.37	PROT
	<b>ATOM</b>	1134 C ASP 367	38.233 4.445 30.574 1.00 14.44	PROT
25	<b>ATOM</b>	1135 O ASP 367	38.457 5.380 29.815 1.00 26.42	PROT
	<b>ATOM</b>	1136 N THR 368	37.619 4.612 31.735 1.00 13.62	PROT
	<b>ATOM</b>	1137 CA THR 368	37.157 5.926 32.160 1.00 13.14	PROT
	<b>ATOM</b>	1138 CB THR 368	36.510 5.853 33.547 1.00 16.53	PROT
	<b>ATOM</b>	1139 OG1 THR 368	35.482 4.856 33.550 1.00 10.44	PROT
30	<b>ATOM</b>	1140 CG2 THR 368	35.928 7.188 33.925 1.00 5.20	PROT
	<b>ATOM</b>	1141 C THR 368	38.291 6.942 32.226 1.00 13.03	PROT
	<b>ATOM</b>	1142 O THR 368	38.114 8.108 31.878 1.00 12.90	PROT
	<b>ATOM</b>	1143 N GLU 369	39.455 6.492 32.686 1.00 9.96	PROT
	<b>ATOM</b>	1144 CA GLU 369	40.616 7.365 32.821 1.00 7.34	PROT
35	<b>ATOM</b>	1145 CB GLU 369	41.673 6.687 33.708 1.00 10.25	PROT
	<b>ATOM</b>	1146 CG GLU 369	41.584 7.113 35.189 1.00 14.56	PROT
	<b>ATOM</b>	1147 CD GLU 369	41.599 5.945 36.167 1.00 19.39	PROT
	<b>ATOM</b>	1148 OE1 GLU 369	42.255 4.922 35.864 1.00 19.65	PROT
	<b>ATOM</b>	1149 OE2 GLU 369	40.954 6.054 37.233 1.00 7.98	PROT
40	<b>ATOM</b>	1150 C GLU 369	41.203 7.768 31.468 1.00 4.33	PROT
	<b>ATOM</b>	1151 O GLU 369	41.467 8.944 31.213 1.00 7.50	PROT
	ATOM	1152 N VAL 370	41.406 6.784 30.603 1.00 12.29	PROT
	<b>ATOM</b>	1153 CA VAL 370	41.927 7.040 29.267 1.00 19.01	PROT
	<b>ATOM</b>	1154 CB VAL 370	42.092 5.726 28.496 1.00 10.10	PROT
45	ATOM	1155 CG1 VAL 370	42.431 6.011 27.049 1.00 8.57	PROT
	ATOM	1156 CG2 VAL 370	43.168 4.877 29.159 1.00 12.40	PROT
	ATOM	1157 C VAL 370	40.896 7.915 28.555 1.00 18.30	PROT
	ATOM	1158 O VAL 370	41.230 8.872 27.855 1.00 17.19	PROT
•	ATOM	1159 N ALA 371	39.633 7.581 28.760 1.00 2.00	PROT
50	ATOM	1160 CA ALA 371	38.549 8.321 28.157 1.00 3.53	PROT

	ATOM	1161 CB ALA 371	37.215 7.728 28.591 1.00 9.17	PROT
	ATOM	1162 C ALA 371	38.603 9.797 28.529 1.00 9.97	PROT
	ATOM	1163 O ALA 371	38.626 10.666 27.655 1.00 24.55	PROT
	ATOM	1164 N LEU 372	38.633 10.082 29.831 1.00 14.85	PROT
5	ATOM	1165 CA LEU 372	38.636 11.463 30.307 1.00 9.24	PROT
	ATOM	1166 CB LEU 372	38.480 11.501 31.830 1.00 8.83	PROT
	ATOM	1167 CG LEU 372	37.043 11.288 32.364 1.00 5.50	PROT
	ATOM	1168 CD1 LEU 372	37.036 10.338 33.553 1.00 2.02	PROT
	ATOM	1169 CD2 LEU 372	36.455 12.626 32.770 1.00 2.00	PROT
10	ATOM	1170 C LEU 372	39.867 12.218 29.870 1.00 10.17	PROT
	ATOM	1171 O LEU 372	39.791 13.413 29.568 1.00 7.23	PROT
	ATOM	1172 N LEU 373	40.996 11.510 29.825 1.00 13.10	PROT
	ATOM	1173 CA LEU 373	42.270 12.078 29.399 1.00 2.00	PROT
	ATOM	1174 CB LEU 373	43.325 10.981 29.381 1.00 2.00	PROT
15	ATOM	1175 CG LEU 373	44.705 11.118 30.045 1.00 9.64	PROT
	ATOM	1176 CD1 LEU 373	44.817 12.382 30.875 1.00 2.00	PROT
	ATOM	1177 CD2 LEU 373	44.955 9.883 30.882 1.00 2.00	PROT
	ATOM	1178 C LEU 373	42.026 12.602 27.987 1.00 6.58	PROT
	ATOM	1179 O LEU 373	42.357 13.738 27.660 1.00 9.73	PROT
20	ATOM	1180 N GLN 374	41.401 11.763 27.165 1.00 9.45	PROT
	ATOM	1181 CA GLN 374	41.076 12.097 25.785 1.00 2.00	PROT
	ATOM	1182 CB GLN 374	40.382 10.914 25.121 1.00 2.00	PROT
	ATOM		41.332 9.896 24.537 1.00 2.00	PROT
	ATOM	1184 CD GLN 374	40.630 8.641 24.095 1.00 2.00	PROT
25	ATOM	1185 OE1 GLN 374	41.261 7.622 23.855 1.00 8.01	PROT
	ATOM	1186 NE2 GLN 374	39.316 8.705 23.989 1.00 2.00	PROT
	ATOM ATOM	1187 C GLN 374	40.187 13.326 25.694 1.00 2.78	PROT
		1188 O GLN 374	40.427 14.213 24.875 1.00 13.91	PROT
30	ATOM ATOM	1189 N ALA 375 1190 CA ALA 375	39.151 13.386 26.521 1.00 2.00	PROT
50	ATOM	1190 CA ALA 373	38.261 14.546 26.505 1.00 2.00 37.128 14.348 27.489 1.00 3.97	PROT
	ATOM	1191 CB ALA 375	39.061 15.801 26.868 1.00 4.60	PROT PROT
	ATOM	1192 C ALA 373 1193 O ALA 375	38.881 16.864 26.274 1.00 8.82	PROT
	ATOM	1194 N VAL 376	39.956 15.667 27.842 1.00 9.01	PROT
35	ATOM	1195 CA VAL 376	40.772 16.790 28.267 1.00 7.36	PROT
23	ATOM	1196 CB VAL 376	41.669 16.401 29.467 1.00 2.30	PROT
	ATOM	1197 CG1 VAL 376	42.597 17.532 29.839 1.00 2.00	PROT
	ATOM	1198 CG2 VAL 376	40.801 16.076 30.646 1.00 9.15	PROT
		1199 C VAL 376	41.629 17.256 27.110 1.00 3.94	PROT
40		1200 O VAL 376	41.788 18.455 26.880 1.00 2.00	PROT
		1201 N LEU 377	42.179 16.297 26.379 1.00 3.92	PROT
	ATOM	1202 CA LEU 377	43.020 16.618 25.239 1.00 5.65	PROT
	ATOM	1203 CB LEU 377	43.714 15.354 24.731 1.00 5.08	PROT
	ATOM	1204 CG LEU 377	45.052 15.005 25.386 1.00 2.00	PROT
45	<b>ATOM</b>	1205 CD1 LEU 377	45.620 13.790 24.719 1.00 2.00	PROT
	ATOM	1206 CD2 LEU 377	46.016 16.157 25.264 1.00 4.14	PROT
	ATOM	1207 C LEU 377	42.173 17.271 24.137 1.00 11.35	PROT
	ATOM	1208 O LEU 377	42.607 18.240 23.515 1.00 8.78	PROT
	ATOM	1209 N LEU 378	40.959 16.766 23.912 1.00 5.62	PROT
50	ATOM	1210 CA LEU 378	40.080 17.352 22.900 1.00 8.57	PROT

	ATOM	1211 CB LEU 378	38.784 16.553 22.788 1.00 5.98	PROT
	ATOM	1212 CG LEU 378	37.847 16.993 21.658 1.00 6.60	PROT
	ATOM	1213 CD1 LEU 378	38.550 16.826 20.329 1.00 2.00	PROT
	ATOM	1214 CD2 LEU 378	36.563 16.172 21.690 1.00 9.27	PROT
5	ATOM	1215 C LEU 378	39.738 18.833 23.146 1.00 10.76	PROT
	<b>ATOM</b>	1216 O LEU 378	40.045 19.689 22.312 1.00 14.81	PROT
	ATOM	1217 N MET 379	39.106 19.139 24.278 1.00 13.15	PROT
	ATOM	1218 CA MET 379	38.735 20.521 24.591 1.00 13.60	PROT
	ATOM	1219 CB MET 379	37.698 20.543 25.709 1.00 12.57	PROT
10	ATOM	1220 CG MET 379	36.425 19.782 25.395 1.00 21.12	PROT
	ATOM	1221 SD MET 379	35.533 20.396 23.927 1.00 15.79	PROT
	ATOM	1222 CE MET 379	34,397 19.099 23.756 1.00 13.95	PROT
	ATOM	1223 C MET 379	39.912 21.419 24.988 1.00 16.01	PROT
	ATOM		39.981 21.897 26.121 1.00 16.95	PROT
15	ATOM	1225 N SER 380	40.824 21.663 24.048 1.00 12.39	PROT
10	ATOM	1226 CA SER 380	41.984 22.506 24.303 1.00 10.77	PROT
	ATOM	1227 CB SER 380	43.248 21.815 23.810 1.00 8.45	PROT
	ATOM	1228 OG SER 380	43.288 20.487 24.286 1.00 17.27	PROT
	ATOM	1229 C SER 380	41.825 23.859 23.621 1.00 15.58	PROT
20	ATOM	1230 O SER 380	42.125 24.019 22.432 1.00 23.09	PROT
20	ATOM	1231 N SER 381	41.368 24.837 24.396 1.00 23.65	PROT
	ATOM	1232 CA SER 381	41.123 26.187 23.904 1.00 25.18	PROT
	ATOM	1232 CR SER 381	40.449 27.018 25.003 1.00 34.78	PROT
	ATOM	1234 OG SER 381	41.250 27.073 26.170 1.00 37.79	PROT
25	ATOM	1235 C SER 381	42.342 26.940 23.388 1.00 19.38	PROT
23	ATOM	1236 O SER 381	42.216 28.032 22.850 1.00 28.81	PROT
	ATOM	1237 N ASP 382	43.519 26.361 23.523 1.00 11.80	PROT
	ATOM	1238 CA ASP 382	44.716 27.057 23.082 1.00 15.78	PROT
	ATOM	1239 CB ASP 382	45.908 26.595 23.909 1.00 33.97	PROT
30	ATOM	1240 CG ASP 382	46.069 25.098 23.891 1.00 48.78	PROT
50	ATOM	1241 OD1 ASP 382	45.169 24.401 24.406 1.00 45.58	PROT
	ATOM	1242 OD2 ASP 382	47.091 24.620 23.356 1.00 56.52	PROT
	ATOM	1243 C ASP 382	45.037 26.888 21.604 1.00 21.28	PROT
	ATOM	1244 O ASP 382	45.907 27.585 21.079 1.00 41.91	PROT
35	ATOM	1245 N ARG 383	44.357 25.971 20.923 1.00 21.81	PROT
33	ATOM	1246 CA ARG 383	44.636 25.773 19.503 1.00 18.95	PROT
	ATOM	1247 CB ARG 383	43.745 24.685 18.921 1.00 8.26	PROT
	ATOM	1248 CG ARG 383	43.580 23.491 19.821 1.00 18.07	PROT
	ATOM	1249 CD ARG 383	44.693 22.487 19.610 1.00 11.10	PROT
40	ATOM	1250 NE ARG 383	44.480 21.261 20.378 1.00 20.54	PROT
70	ATOM	1250 NE ARG 383	45.460 20.462 20.786 1.00 18.25	PROT
	ATOM	1251 CZ ARG 383	45.187 19.365 21.481 1.00 5.24	PROT
	ATOM	1252 NH1 ARG 383	46.717 20.765 20.495 1.00 19.21	PROT
	ATOM	1254 C ARG 383	44.420 27.064 18.728 1.00 19.64	PROT
45	ATOM	1255 O ARG 383	43.493 27.828 19.001 1.00 17.46	PROT
43	ATOM	1256 N PRO 384	45.298 27.342 17.762 1.00 25.37	PROT
	ATOM	1256 N PRO 384 1257 CD PRO 384	46.485 26.567 17.359 1.00 25.37	PROT
	ATOM	1257 CD PRO 384 1258 CA PRO 384	45.124 28.569 16.983 1.00 27.53	PROT
		1258 CA PRO 384 1259 CB PRO 384	45.124 28.369 10.383 1.00 27.33 46.422 28.693 16.181 1.00 18.75	PROT
50	ATOM ATOM		47.041 27.338 16.190 1.00 27.78	PROT
50	ATOM	1260 CG PRO 384	4/.041 2/./0 10.170 1.00 2/./8	1 1/01

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WO 99/26966

PCT/US98/25296

	ATOM	1261 C PRO 384	43.895 28.476 16.081 1.00 28.76	PROT
	ATOM	1262 O PRO 384	43.562 27.402 15.560 1.00 31.18	PROT
	ATOM	1263 N GLY 385	43.215 29.606 15.917 1.00 27.37	PROT
	ATOM	1264 CA GLY 385	42.039 29.638 15.073 1.00 26.98	PROT
5	ATOM	1265 C GLY 385	40.728 29.442 15.803 1.00 27.46	PROT
	ATOM	1266 O GLY 385	39.689 29.911 15.339 1.00 31.99	PROT
	ATOM	1267 N LEU 386	40.756 28.756 16.939 1.00 34.99	PROT
	ATOM	1268 CA LEU 386	39.524 28.515 17.673 1.00 37.24	PROT
	ATOM	1269 CB LEU 386	39.820 27.947 19.059 1.00 26.60	PROT
10	ATOM	1270 CG LEU 386	40.233 26.472 18.988 1.00 32.45	PROT
	ATOM	1271 CD1 LEU 386	40.177 25.859 20.363 1.00 34.82	PROT
	ATOM	1272 CD2 LEU 386	39.314 25.719 18.030 1.00 29.64	PROT
	ATOM	1273 C LEU 386	38.733 29.795 17.778 1.00 36.93	PROT
	ATOM	1274 O LEU 386	39.291 30.881 17.674 1.00 37.60	PROT
15	ATOM	1275 N ALA 387	37.427 29.665 17.962 1.00 31.47	PROT
	ATOM	1276 CA ALA 387	36.578 30.832 18.058 1.00 28.80	PROT
	ATOM	1277 CB ALA 387	35.553 30.814 16.950 1.00 41.01	PROT
	ATOM	1278 C ALA 387	35.890 30.864 19.400 1.00 28.89	PROT
	ATOM	1279 O ALA 387	35.998 31.842 20.133 1.00 30.62	PROT
20	ATOM	1280 N CYS 388	35.167 -29.797 19.710 1.00 25.92	PROT
	ATOM	1281 CA CYS 388	34.469 29.712 20.978 1.00 26.90	PROT
	ATOM	1282 CB CYS 388	33.224 28.823 20.826 1.00 21.38	PROT
	ATOM	1283 SG CYS 388	31.625 29.732 20.698 1.00 33.66	PROT
	ATOM	1284 C CYS 388	35.443 29.159 22.040 1.00 31.18	PROT
25	ATOM	1285 O CYS 388	35.272 28.054 22.552 1.00 36.57	PROT
	ATOM	1286 N VAL 389	36.473 29.951 22.346 1.00 20.22	PROT
	ATOM	1287 CA VAL 389	37.511 29.622 23.327 1.00 16.02	PROT
	ATOM	1288 CB VAL 389	38.554 30.737 23.381 1.00 9.80	PROT
	ATOM	1289 CG1 VAL 389	39.526 30.480 24.498 1.00 16.03	PROT
30	ATOM	1290 CG2 VAL 389	39.257 30.843 22.056 1.00 16.27	PROT
	ATOM	1291 C VAL 389	36.977 29.425 24.753 1.00 18.85	PROT
	ATOM	1292 O VAL 389	37.066 28.336 25.323 1.00 24.21	PROT
	ATOM	1293 N GLU 390	36.461 30.500 25.337 1.00 5.06	PROT
	ATOM	1294 CA GLU 390	35.908 30.434 26.660 1.00 2.00	PROT
35	ATOM	1295 CB GLU 390	35.092 31.684 26.952 1.00 5.13	PROT
	ATOM	1296 C GLU 390	35.047 29.184 26.817 1.00 3.75	PROT
	ATOM	1297 O GLU 390	35.252 28.419 27.754 1.00 23.35	PROT
	ATOM	1298 N ARG 391	34.103 28.938 25.915 1.00 14.06	PROT
	ATOM	1299 CA ARG 391	33.248 27.754 26.093 1.00 26.18	PROT
40	ATOM	1300 CB ARG 391	32.121 27.699 25.049 1.00 31.84	PROT
	ATOM	1301 CG ARG 391	30.843 27.040 25.601 1.00 47.73	PROT
	ATOM	1302 CD ARG 391	29.882 26.572 24.512 1.00 58.24	PROT
	ATOM	1303 NE ARG 391	29.879 27.487 23.378 1.00 66.80	PROT
	ATOM	1304 CZ ARG 391	29.001 28.470 23.211 1.00 69.56	PROT
45	ATOM	1305 NH1 ARG 391	29.088 29.255 22.139 1.00 66.99	PROT
	ATOM	1306 NH2 ARG 391	28.034 28.663 24.105 1.00 56.08	PROT
	ATOM	1307 C ARG 391	33.979 26.415 26.110 1.00 23.65	PROT
	ATOM	1308 O ARG 391	33.561 25.479 26.794 1.00 28.58	PROT
	ATOM	1309 N ILE 392		PROT
50	ATOM	1310 CA ILE 392	35.812 25.077 25.335 1.00 19.03	PROT

	ATOM	1311 CB ILE 392	36.804 25.063 24.165 1.00 22.30	PROT
	ATOM	1312 CG2 ILE 392	37.971 24.130 24.467 1.00 21.71	PROT
	ATOM	1313 CG1 ILE 392	36.074 24.614 22.892 1.00 23.47	PROT
	ATOM	1314 CD1 ILE 392	36.245 25.551 21.707 1.00 4.13	PROT
5	ATOM	1315 C ILE 392	36.544 24.907 26.671 1.00 25.03	PROT
	ATOM	1316 O ILE 392	36.728 23.783 27.153 1.00 26.11	PROT
	ATOM	1317 N GLU 393	36.947 26.029 27.266 1.00 30.74	PROT
	ATOM	1318 CA GLU 393	37.630 26.021 28.558 1.00 23.39	PROT
	ATOM	1319 CB GLU 393	38.073 27.430 28.930 1.00 27.18	PROT
10	ATOM	1320 CG GLU 393	39.435 27.817 28.402 1.00 41.39	PROT
	ATOM	1321 CD GLU 393	39.990 29.051 29.093 1.00 47.72	PROT
	ATOM	1322 OE1 GLU 393	39.365 29.524 30.070 1.00 39.94	PROT
	ATOM	1323 OE2 GLU 393	41.051 29.547 28.653 1.00 51.17	PROT
	ATOM	1324 C GLU 393	36.655 25.516 29.610 1.00 21.72	PROT
15	ATOM	1325 O GLU 393	36.942 24.574 30.344 1.00 22.82	PROT
	ATOM	1326 N LYS 394	35.497 26.163 29.676 1.00 9.64	PROT
	ATOM	1327 CA LYS 394	34.462 25.779 30.618 1.00 11.56	PROT
	ATOM	1328 CB LYS 394	33.177 26,557 30.338 1.00 7.52	PROT
	ATOM	1329 C LYS 394	34.213 24.280 30.492 1.00 16.31	PROT
20	ATOM	1330 O LYS 394	34.000 23.594 31.498 1.00 24.52	PROT
	<b>ATOM</b>	1331 N TYR 395	34.251 23.763 29.264 1.00 12.79	PROT
	<b>ATOM</b>	1332 CA TYR 395	34.033 22.332 29.057 1.00 19.02	PROT
	ATOM	1333 CB TYR 395	33.803 22.025 27.572 1.00 27.90	PROT
	ATOM	1334 CG TYR 395	32.454 22.456 27.027 1.00 31.64	PROT
25	<b>ATOM</b>	1335 CD1 TYR 395	32.136 22.267 25.684 1.00 30.15	PROT
	ATOM	1336 CE1 TYR 395	30.927 22.695 25.160 1.00 28.34	PROT
	ATOM	1337 CD2 TYR 395	31.514 23.085 27.835 1.00 34.21	PROT
	ATOM	1338 CE2 TYR 395	30.298 23.518 27.317 1.00 34.01	PROT
	ATOM	1339 CZ TYR 395	30.014 23.322 25.979 1.00 33.73	PROT
30	ATOM	1340 OH TYR 395	28.824 23.785 25.453 1.00 44.99	PROT
	ATOM	1341 C TYR 395	35.208 21.490 29.584 1.00 19.03	PROT
	ATOM	1342 O TYR 395	35.003 20.494 30.277 1.00 25.23	PROT
	ATOM	1343 N GLN 396	36.437 21.883 29.256 1.00 17.76	PROT
	ATOM	1344 CA GLN 396	37.596 21.134 29.725 1.00 13.73	PROT
35	ATOM	1345 CB GLN 396	38.905 21.766 29.240 1.00 2.45	PROT
	ATOM	1346 CG GLN 396	40.061 20.767 29.110 1.00 2.00	PROT
	ATOM	1347 CD GLN 396	41.388 21.439 28.799 1.00 5.12	PROT
	ATOM	1348 OE1 GLN 396	41.706 22.484 29.359 1.00 10.11	PROT
	ATOM	1349 NE2 GLN 396	42.169 20.840 27.903 1.00 9.09	PROT
40		1350 C GLN 396	37.562 21.149 31.238 1.00 17.65	PROT
	ATOM	1351 O GLN 396	37.802 20.125 31.894 1.00 9.63	PROT
	ATOM	1352 N ASP 397	37.250 22.319 31.787 1.00 6.69	PROT
	ATOM		37.178 22.476 33.226 1.00 9.36	PROT
	ATOM		36.732 23.893 33.570 1.00 11.44	PROT
45	ATOM	1355 CG ASP 397	37.867 24.891 33.446 1.00 18.32	PROT
	ATOM		39.033 24.438 33.397 1.00 24.00	PROT
	ATOM		37.615 26.114 33.395 1.00 20.67	PROT
	ATOM		36.215 21.443 33.771 1.00 7.77	PROT
	ATOM		36.497 20.771 34.761 1.00 7.66	PROT
50	ATOM	1360 N SER 398	35.087 21.293 33.093 1.00 9.19	PROT

	АТОМ	1361 CA SER 398	34.094 20.322 33.508 1.00 14.18	PROT
	ATOM	1362 CB SER 398	32.916 20.334 32.542 1.00 12.11	PROT
	ATOM	1363 OG SER 398	32.406 21.650 32.423 1.00 31.95	PROT
	ATOM	1364 C SER 398	34.712 18.939 33.556 1.00 11.47	PROT
5	ATOM	1365 O SER 398	34.591 18.227 34.551 1.00 21.11	PROT
•	ATOM	1366 N PHE 399	35.394 18.565 32.485 1.00 18.68	PROT
	ATOM	1367 CA PHE 399	36.017 17.252 32.417 1.00 24.93	PROT
	ATOM	1368 CB PHE 399	36.587 17.012 31.014 1.00 23.38	PROT
	ATOM	1369 CG PHE 399	35,543 16,705 29,981 1.00 20.19	PROT
10	ATOM	1370 CD1 PHE 399	35.224 17.638 28.997 1.00 22.94	PROT
- •	ATOM	1371 CD2 PHE 399	34.878 15.486 29.988 1.00 8.62	PROT
	ATOM	1372 CE1 PHE 399	34.257 17.361 28.029 1.00 12.53	PROT
	ATOM	1373 CE2 PHE 399	33.914 15.201 29.027 1.00 19.25	PROT
	ATOM	1374 CZ PHE 399	33.604 16.143 28.044 1.00 15.15	PROT
15	ATOM	1375 C PHE 399	37.113 17.097 33.463 1.00 23.06	PROT
	ATOM	1376 O PHE 399	37.210 16.063 34.137 1.00 15.58	PROT
	ATOM	1377 N LEU 400	37.932 18.131 33.604 1.00 22.12	PROT
	ATOM	1378 CA LEU 400	39.017 18.095 34.567 1.00 18.27	PROT
	ATOM	1379 CB LEU 400	39.846 19.372 34.461 1.00 10.06	PROT
20	ATOM	1380 CG LEU 400	41.021 19.248 33.491 1.00 8.13	PROT
	<b>ATOM</b>	1381 CD1 LEU 400	41.616 20.594 33.195 1.00 2.00	PROT
	ATOM	1382 CD2 LEU 400	42.055 18.333 34.095 1.00 13.73	PROT
	ATOM	1383 C LEU 400	38.527 17.892 36.002 1.00 24.79	PROT
	<b>ATOM</b>	1384 O LEU 400	39.189 17.228 36.787 1.00 26.46	PROT
25	<b>ATOM</b>	1385 N LEU 401	37.371 18.447 36.354 1.00 21.93	PROT
	ATOM	1386 CA LEU 401	36.862 18.268 37.707 1.00 17.21	PROT
	ATOM	1387 CB LEU 401	35.766 19.285 38.022 1.00 19.27	PROT
	ATOM	1388 CG LEU 401	35.538 19.547 39.515 1.00 16.76	PROT
	ATOM	1389 CD1 LEU 401	36.652 20.403 40.085 1.00 2.00	PROT
30	ATOM	1390 CD2 LEU 401	34.206 20.235 39.687 1.00 14.41	PROT
	ATOM	1391 C LEU 401	36.316 16.864 37.879 1.00 18.03	PROT
	ATOM	1392 O LEU 401	36.482 16.250 38.925 1.00 28.63	PROT
	ATOM	1393 N ALA 402	35.656 16.346 36.856 1.00 9.30	PROT
	ATOM	1394 CA ALA 402	35.124 15.000 36.951 1.00 7.03	PROT
35	ATOM	1395 CB ALA 402	34.233 14.703 35.758 1.00 14.15	PROT
	ATOM	1396 C ALA 402	36.298 14.029 36.989 1.00 7.68	PROT
	ATOM	1397 O ALA 402	36.294 13.054 37.739 1.00 2.00	PROT
	ATOM	1398 N PHE 403	37.311 14.305 36.178 1.00 4.49	PROT
40	ATOM	1399 CA PHE 403	38.477 13.439 36.140 1.00 9.18	PROT
40	ATOM	1400 CB PHE 403	39.510 13.977 35.138 1.00 12.80	PROT
	ATOM	1401 CG PHE 403	40.545 12.957 34.693 1.00 5.42	PROT
	ATOM	1402 CD1 PHE 403	41.590 13.334 33.859 1.00 2.00	PROT
	ATOM	1403 CD2 PHE 403	40.480 11.634 35.103 1.00 2.00	PROT
4.5	ATOM	1404 CE1 PHE 403	42.546 12.410 33.448 1.00 2.00	PROT
45	ATOM	1405 CE2 PHE 403	41.440 10.711 34.688 1.00 2.00	PROT
	ATOM	1406 CZ PHE 403	42.468 11.100 33.863 1.00 2.00	PROT
	ATOM	1407 C PHE 403	39.080 13.366 37.539 1.00 10.08	PROT
	ATOM	1408 O PHE 403	39.207 12.279 38.097 1.00 8.23	PROT
50	ATOM	1409 N GLU 404	39.451 14.514 38.103 1.00 12.64	PROT
50	ATOM	1410 CA GLU 404	40.030 14.546 39.448 1.00 19.23	PROT

ATOM 1413 CD GLU 404 ATOM 1414 OEI GLU 404 ATOM 1416 C GLU 404 ATOM 1417 O GLU 404 ATOM 1418 N HIS 405 ATOM 1419 CA HIS 405 ATOM 1419 CA HIS 405 ATOM 1420 CB HIS 405 ATOM 1420 CB HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1422 ND1 HIS 405 ATOM 1424 CBI HIS 405 ATOM 1425 NE2 HIS 405 ATOM 1426 C HIS 405 ATOM 1427 O HIS 405 ATOM 1428 N TYR 406 ATOM 1430 CB TYR 406 ATOM 1440 CD		ATOM ATOM	1411 CB GLU 404 40.227 15.989 39.942 1.00 19.80 PROT 1412 CG GLU 404 41.532 16.220 40.728 1.00 24.03 PROT
ATOM 1414 OE1 GLU 404 ATOM 1415 OE2 GLU 404 ATOM 1416 C GLU 404 ATOM 1417 O GLU 404 ATOM 1417 O GLU 404 ATOM 1418 N HIS 405 ATOM 1418 N HIS 405 ATOM 1419 CA HIS 405 ATOM 1419 CA HIS 405 ATOM 1420 CB HIS 405 ATOM 1421 CG HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1423 ND1 HIS 405 ATOM 1424 CEI HIS 405 ATOM 1426 C HIS 405 ATOM 1426 C HIS 405 ATOM 1427 O HIS 405 ATOM 1428 N TYR 406 ATOM 1429 CA TYR 406 ATOM 1431 CG TYR 406 ATOM 1431 CG TYR 406 ATOM 1432 CD1 TYR 406 ATOM 1434 CD2 TYR 406 ATOM 1435 CE2 TYR 406 ATOM 1436 CZ TYR 406 ATOM 1437 OH TYR 406 ATOM 1437 OH TYR 406 ATOM 1443 CG2 ILE 407 ATOM 1443 CG2 ILE 407 ATOM 1444 CG ILE 407 ATOM 1444 CG ILE 407 ATOM 1445 CD ASN 408 ATOM 1			
ATOM			
ATOM 1416 C GLU 404 ATOM 1417 O GLU 404 ATOM 1418 N HIS 405 ATOM 1418 N HIS 405 ATOM 1419 CA HIS 405 ATOM 1420 CB HIS 405 ATOM 1421 CG HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1423 ND1 HIS 405 ATOM 1424 CEI HIS 405 ATOM 1426 C HIS 405 ATOM 1426 C HIS 405 ATOM 1427 O HIS 405 ATOM 1429 CA TYR 406 ATOM 1430 CB TYR 406 ATOM 1431 CG TYR 406 ATOM 1431 CG TYR 406 ATOM 1432 CD1 TYR 406 ATOM 1433 CEI TYR 406 ATOM 1435 CEZ TYR 406 ATOM 1436 CZ TYR 406 ATOM 1437 OH TYR 406 ATOM 1438 C TYR 406 ATOM 1438 C TYR 406 ATOM 1439 O TYR 406 ATOM 1430 CB TIR 406 ATOM 1431 CG IIE 407 ATOM 1431 CG IIE 407 ATOM 1432 CD1 TYR 406 ATOM 1433 CEI TYR 406 ATOM 1434 CD2 TYR 406 ATOM 1436 CZ TYR 406 ATOM 1437 OH TYR 406 ATOM 1438 C TYR 406 ATOM 1444 CGI ILE 407 ATOM 1444 CGI ILE 407 ATOM 1444 CGI ILE 407 ATOM 1445 CD1 ILE 407 ATOM 1445 CD1 ILE 407 ATOM 1445 CD IIE 407 ATOM 1446 CR IIE 407 ATOM 1447 C A SNN 408 ATOM 1445 CD II SN	5		
ATOM 1418 N HIS 405 ATOM 1419 CA HIS 405 ATOM 1420 CB HIS 405 ATOM 1421 CG HIS 405 ATOM 1421 CG HIS 405 ATOM 1422 CD2 HIS 405 ATOM 1423 ND1 HIS 405 ATOM 1424 CEI HIS 405 ATOM 1425 NE2 HIS 405 ATOM 1426 C HIS 405 ATOM 1427 O HIS 405 ATOM 1427 O HIS 405 ATOM 1428 N TYR 406 ATOM 1429 CA TYR 406 ATOM 1430 CB TYR 406 ATOM 1431 CG TYR 406 ATOM 1431 CD TYR 406 ATOM 1432 CD1 TYR 406 ATOM 1433 CEI TYR 406 ATOM 1436 CZ TYR 406 ATOM 1436 CZ TYR 406 ATOM 1437 O HIS 405 ATOM 1438 C TYR 406 ATOM 1438 C TYR 406 ATOM 1436 CZ TYR 406 ATOM 1437 O HIS 405 ATOM 1438 C TYR 406 ATOM 1438 C TYR 406 ATOM 1439 O TYR 406 ATOM 1440 N ILE 407 ATOM 1441 CA ILE 407 ATOM 1442 CB ILE 407 ATOM 1443 CG2 ILE 407 ATOM 1444 CGI ILE 407 ATOM 1445 CD1 ILE 407 ATOM 1446 C ILE 407 ATOM 1447 O ILE 407 ATOM 1447 O ILE 407 ATOM 1447 O ILE 407 ATOM 1446 C ILE 407 ATOM 1447 O ILE 407 ATOM 1447 O ILE 407 ATOM 1447 CA SN 408 ATOM 1447 CA SN 408 ATOM 1445 CD1 SN 408 ATOM 1445 CD1 SN 408 ATOM 1445 CD SAN 408 ATOM 1445 C CHANN 408 ATOM 1446 C CHANN 408 ATOM 1446 C CHANN 408 ATOM 1447 O HILE 407 ATOM 1446 C CHANN 408 ATOM 1445 C CHANN 408 ATOM 1455 O ASN 408 ATOM 1456 O TYR 409 ATOM 1457 C A TYR 409 ATOM 1458 C CHANN 409 ATOM 1458 C CHANN 408 ATOM 1458 C CHANN 408 ATOM 1459 C CHANN 408 ATOM 1458 C CHANN 408 ATOM 1458 C CHANN 408 AT	,		
ATOM 1418 N HIS 405 37.815 14.108 40.358 1.00 10.26 PROT ATOM 1419 CA HIS 405 36.870 13.446 41.240 1.00 7.78 PROT ATOM 1421 CG HIS 405 35.312 15.393 41.630 1.00 15.49 PROT ATOM 1422 CD2 HIS 405 36.223 16.260 42.134 1.00 17.97 PROT ATOM 1423 ND1 HIS 405 34.265 17.242 42.210 1.00 27.50 PROT ATOM 1424 CE1 HIS 405 34.265 17.242 42.210 1.00 27.50 PROT ATOM 1425 NE2 HIS 405 36.856 11.936 41.694 1.00 21.57 PROT ATOM 1426 C HIS 405 36.856 11.936 41.694 1.00 21.57 PROT ATOM 1426 C HIS 405 36.856 11.936 41.095 1.00 14.88 PROT ATOM 1427 O HIS 405 36.641 11.155 41.935 1.00 22.11 PROT ATOM 1428 N TYR 406 37.091 11.512 39.767 1.00 16.52 PROT ATOM 1429 CA TYR 406 37.097 9.808 37.989 1.00 9.90 PROT ATOM 1431 CG TYR 406 37.007 9.808 37.989 1.00 9.90 PROT ATOM 1432 CD1 TYR 406 37.801 6.204 37.041 1.00 13.48 PROT ATOM 1434 CD2 TYR 406 36.840 8.346 37.657 1.00 2.00 PROT ATOM 1435 CE2 TYR 406 37.801 6.204 37.044 1.00 13.48 PROT ATOM 1436 CZ TYR 406 36.481 4.287 36.804 1.00 2.00 PROT ATOM 1437 OH TYR 406 36.481 4.287 36.804 1.00 2.00 PROT ATOM 1438 C TYR 406 36.481 4.287 36.804 1.00 2.00 PROT ATOM 1439 O TYR 406 38.340 9.466 40.071 1.00 9.54 PROT ATOM 1441 CA ILE 407 ATOM 1442 CB ILE 407 40.638 1.00 1.01 1.49 PROT ATOM 1444 CGI ILE 407 40.638 1.00 1.00 11.28 PROT ATOM 1444 CGI ILE 407 40.438 9.426 42.091 1.00 11.49 PROT ATOM 1444 CGI ILE 407 40.438 9.426 42.091 1.00 11.44 PROT ATOM 1444 CGI ILE 407 40.438 9.426 42.091 1.00 11.44 PROT ATOM 1445 CD3 ASN 408 39.649 12.840 44.657 1.00 3.57 PROT ATOM 1445 CD3 ASN 408 39.649 12.840 44.657 1.00 2.00 PROT ATOM 1445 CD3 ASN 408 39.649 12.840 44.657 1.00 2.00 PROT ATOM 1450 CB ASN 408 39.649 12.840 44.657 1.00 2.00 PROT ATOM 1455 O ASN 408 38.868 9.078 44.432 1.00 6.49 PROT ATOM 1456 N TYR 409 36.900 7.816 4389 42.668 1.00 12.55 PROT ATOM 1455 O ASN 408 39.489 12.840 44.657 1.00 2.00 PROT ATOM 1455 O ASN 408 39.489 12.840 44.657 1.00 2.00 PROT ATOM 1455 O ASN 408 39.282 8.187 43.766 1.00 2.00 PROT ATOM 1455 O ASN 408 39.489 12.840 44.657 1.00 2.00 PROT ATOM 1455 O ASN 408 39.282 8.188 44.656			
ATOM   1419   CA HIS   405   36.870   13.446   41.240   1.00   7.78   PROT   ATOM   1421   CG HIS   405   35.473   14.023   41.054   1.00   3.47   PROT   ATOM   1422   CD2 HIS   405   35.312   15.393   41.630   1.00   15.49   PROT   ATOM   1423   ND1 HIS   405   34.096   16.036   41.694   1.00   21.57   PROT   ATOM   1424   CEI HIS   405   34.265   17.242   42.210   1.00   27.50   PROT   ATOM   1425   NE2 HIS   405   34.265   17.242   42.210   1.00   27.50   PROT   ATOM   1426   C   HIS   405   36.841   11.155   41.935   1.00   22.11   PROT   ATOM   1427   C   HIS   405   36.846   11.936   41.095   1.00   14.88   PROT   ATOM   1428   N   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1430   CB   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1431   CG   TYR   406   35.487   36.384   37.657   1.00   2.00   PROT   ATOM   1433   CEI   TYR   406   35.433   6.382   37.338   1.00   8.78   PROT   ATOM   1435   CEZ   TYR   406   35.433   6.382   37.338   1.00   15.34   PROT   ATOM   1436   CZ   TYR   406   36.441   4.287   36.864   1.00   2.00   PROT   ATOM   1436   CZ   TYR   406   36.441   4.287   36.864   1.00   2.00   PROT   ATOM   1430   C TYR   406   36.548   5.624   37.073   1.00   15.64   PROT   ATOM   1436   C TYR   406   36.431   4.287   36.864   1.00   2.00   PROT   ATOM   1440   N   ILE   407   ATOM   1441   CA   ILE   407   ATOM   1442   CB   ILE   407   ATOM   1444   CB   ILE   407   ATOM   1445   CB   ILE   407   ATOM   1445   CD   ILE   407   ATOM   1445   CB   ASN   408   38.758   11.535   44.629   1.00   13.44   PROT   ATOM   1445   CB   ASN   408   38.758   13.949   44.6657   1.00   3.57   PROT   ATOM   1445   CB   ASN   408   38.758   13.949   44.6657   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.6657   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.6657   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.6657   1.00   2.00   PROT   ATOM   1455   O ASN   408   38.758   13.949   4			
ATOM   1420   CB   HIS   405   ATOM   1421   CG   HIS   405   ATOM   1422   CD2   HIS   405   ATOM   1422   CD2   HIS   405   35.312   15.393   41.630   1.00   15.49   PROT   ATOM   1424   CE1   HIS   405   34.096   16.036   41.694   1.00   21.57   PROT   ATOM   1425   CE1   HIS   405   34.265   17.242   42.210   1.00   27.50   PROT   ATOM   1426   C   HIS   405   35.547   17.403   42.485   1.00   13.53   PROT   ATOM   1427   O   HIS   405   ATOM   1428   N   TYR   406   ATOM   1429   CA   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1430   CB   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1431   CG   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1432   CD1   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1433   CE1   TYR   406   37.085   10.083   39.491   1.00   14.35   PROT   ATOM   1434   CD2   TYR   406   35.433   6.382   37.386   1.00   8.78   PROT   ATOM   1435   CE2   TYR   406   37.835   37.386   1.00   8.78   PROT   ATOM   1436   CZ   TYR   406   37.831   6.204   37.041   10.01   3.48   PROT   ATOM   1437   OH   TYR   406   36.431   4.287   36.804   1.00   2.00   PROT   ATOM   1440   N   ILE   407   ATOM   1441   CA   ILE   407   ATOM   1442   CB   ILE   407   ATOM   1444   CB   ILE   407   ATOM   1444   CB   ILE   407   ATOM   1445   CD1   ILE   407   ATOM   1446   CD1   ILE   407   ATOM   1447   O   ILE   407   ATOM   1448   N   ASN   408   39.493   1.091   1.00   1.28   PROT   ATOM   1445   CD1   ATOM   39.933   10.488   42.792   1.00   1.34   PROT   ATOM   1445   CD1   ASN   408   38.758   13.949   44.669   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.669   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.669   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.669   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.669   1.00   2.00   PROT   ATOM   1450   CB   ASN   408   38.758   13.949   44.669   1.00   2.00			
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40       ATOM       1450       CB       ASN       408       38.758       11.535       44.629       1.00       2.00       PROT         ATOM       1451       CG       ASN       408       39.499       12.840       44.657       1.00       3.57       PROT         ATOM       1452       OD1       ASN       408       40.733       12.859       44.656       1.00       14.35       PROT         ATOM       1453       ND2       ASN       408       38.758       13.949       44.689       1.00       2.00       PROT         ATOM       1454       C       ASN       408       38.868       9.078       44.432       1.00       6.49       PROT         45       ATOM       1455       O       ASN       408       39.282       8.187       45.178       1.00       10.45       PROT         ATOM       1456       N       TYR       409       36.900       7.816       43.893       1.00       9.20       PROT         ATOM       1458       CB       TYR       409       35.879       7.783       42.760       1.00       11.66       PROT         ATOM       1459       CG			
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ATOM 1454 C ASN 408 38.868 9.078 44.432 1.00 6.49 PROT  45 ATOM 1455 O ASN 408 39.282 8.187 45.178 1.00 10.45 PROT  ATOM 1456 N TYR 409 37.731 8.987 43.766 1.00 2.00 PROT  ATOM 1457 CA TYR 409 36.900 7.816 43.893 1.00 9.20 PROT  ATOM 1458 CB TYR 409 35.879 7.783 42.760 1.00 11.66 PROT  ATOM 1459 CG TYR 409 35.121 6.489 42.683 1.00 12.54 PROT			
45 ATOM 1455 O ASN 408 39.282 8.187 45.178 1.00 10.45 PROT ATOM 1456 N TYR 409 37.731 8.987 43.766 1.00 2.00 PROT ATOM 1457 CA TYR 409 36.900 7.816 43.893 1.00 9.20 PROT ATOM 1458 CB TYR 409 35.879 7.783 42.760 1.00 11.66 PROT ATOM 1459 CG TYR 409 35.121 6.489 42.683 1.00 12.54 PROT		<b>ATOM</b>	1453 ND2 ASN 408 38.758 13.949 44.689 1.00 2.00 PROT
ATOM 1456 N TYR 409 37.731 8.987 43.766 1.00 2.00 PROT ATOM 1457 CA TYR 409 36.900 7.816 43.893 1.00 9.20 PROT ATOM 1458 CB TYR 409 35.879 7.783 42.760 1.00 11.66 PROT ATOM 1459 CG TYR 409 35.121 6.489 42.683 1.00 12.54 PROT		ATOM	1454 C ASN 408 38.868 9.078 44.432 1.00 6.49 PROT
ATOM 1457 CA TYR 409 36.900 7.816 43.893 1.00 9.20 PROT ATOM 1458 CB TYR 409 35.879 7.783 42.760 1.00 11.66 PROT ATOM 1459 CG TYR 409 35.121 6.489 42.683 1.00 12.54 PROT	45	<b>ATOM</b>	1455 O ASN 408 39.282 8.187 45.178 1.00 10.45 PROT
ATOM 1458 CB TYR 409 35.879 7.783 42.760 1.00 11.66 PROT ATOM 1459 CG TYR 409 35.121 6.489 42.683 1.00 12.54 PROT		<b>ATOM</b>	1456 N TYR 409 37.731 8.987 43.766 1.00 2.00 PROT
ATOM 1459 CG TYR 409 35.121 6.489 42.683 1.00 12.54 PROT			
50 ATOM 1460 CD1 TYR 409 33.984 6.281 43.456 1.00 29.23 PROT			
	50	АТОМ	1460 CD1 TYR 409 33.984 6.281 43.456 1.00 29.23 PROT

	1 TO 1 1	1461 OF1 TVD 400	22 205 5 277 42 402 1 00 25 45	DDOT
	ATOM	1461 CE1 TYR 409	33.285 5.077 43.403 1.00 25.45	PROT
	ATOM	1462 CD2 TYR 409	35.547 5.465 41.850 1.00 24.96	PROT
	ATOM	1463 CE2 TYR 409	34.860 4.259 41.788 1.00 33.40	PROT
_	ATOM	1464 CZ TYR 409	33.733 4.074 42.567 1.00 24.27	PROT
5	ATOM	1465 OH TYR 409	33.065 2.883 42.509 1.00 32.72	PROT
	ATOM		37.753 6.553 43.867 1.00 13.96	PROT
	ATOM		37.730 5.763 44.804 1.00 29.48	PROT
	ATOM	1468 N ARG 410	38.531 6.399 42.803 1.00 23.04	PROT
10	ATOM	1469 CA ARG 410	39.377 5.230 42.588 1.00 22.09	PROT
10	ATOM	1470 CB ARG 410	39.982 5.327 41.190 1.00 13.24	PROT
	ATOM	1471 CG ARG 410	38.947 5.399 40.090 1.00 14.01	PROT
	ATOM	1472 CD ARG 410	38.934 4.111 39.275 1.00 16.49	PROT
	ATOM	1473 NE ARG 410	40.227 3.848 38.651 1.00 9.77	PROT
	ATOM	1474 CZ ARG 410	40.617 2.651 38.239 1.00 11.38	PROT
15	ATOM	1475 NH1 ARG 410	41.806 2.493 37.685 1.00 14.94	PROT
	ATOM	1476 NH2 ARG 410	39.810 1.613 38.375 1.00 12.78	PROT
	ATOM		40.486 4.914 43.604 1.00 24.49	PROT
	ATOM		40.860 3.753 43.780 1.00 12.85	PROT
20	ATOM		41.023 5.931 44.262 1.00 24.16	PROT
20	ATOM	1480 CA LYS 411	42.085 5.706 45.235 1.00 27.14	PROT
	ATOM	1481 CB LYS 411	41.525 5.069 46.516 1.00 37.40	PROT
	ATOM	1482 CG LYS 411	40.317 5.779 47.103 1.00 35.00	PROT
	ATOM	1483 CD LYS 411	39.406 4.788 47.804 1.00 40.83	PROT
25	ATOM	1484 CE LYS 411	38.414 5.496 48.725 1.00 58.04	PROT
25	ATOM	1485 NZ LYS 411	38.833 5.496 50.168 1.00 54.40	PROT
	ATOM		43.186 4.814 44.664 1.00 28.02	PROT
	ATOM		43.209 3.598 44.876 1.00 25.00	PROT
	ATOM			PROT
20	ATOM		45.223 4.738 43.332 1.00 26.70	PROT
30	ATOM		45.756 5.491 42.104 1.00 29.28	PROT
	ATOM		44.953 5.289 40.857 1.00 18.44	PROT PROT
	ATOM	1492 CD2 HIS 412	43.783 5.836 40.451 1.00 19.98	
	ATOM	1493 ND1 HIS 412	45.366 4.465 39.833 1.00 16.33	PROT
25	ATOM	1494 CE1 HIS 412	44.486 4.513 38.850 1.00 24.80	PROT
35	ATOM	1495 NE2 HIS 412	43.516 5.338 39.200 1.00 23.01 6.281 4.788 44.406 1.00 20.73	PROT
	ATOM			PROT
	ATOM			PROT
	ATOM			PROT
40	ATOM		48.183 3.788 45.465 1.00 28.09	PROT
40	ATOM		48.219 2.426 46.144 1.00 21.71	PROT
	ATOM		46.906 2.053 46.759 1.00 44.26	PROT
	ATOM	1502 CD2 HIS 413	46.140 0.941 46.632 1.00 43.48	PROT
	ATOM	1503 ND1 HIS 413	46.214 2.902 47.600 1.00 40.00	PROT
4.5	ATOM		45.080 2.328 47.962 1.00 47.35	PROT
45	ATOM	1505 NE2 HIS 413	45.011 1.137 47.390 1.00 35.50	PROT
	ATOM			PROT
	ATOM			PROT
	ATOM		49.555 5.439 44.411 1.00 18.32	PROT
50	ATOM	1509 CA VAL 414	50.726 6.069 43.820 1.00 22.60	PROT
50	ATOM	1510 CB VAL 414	50.718 5.966 42.290 1.00 32.50	PROT

	ATOM	1511 CGI VAL 414 51.63	6 7.026 41.694 1.00 33.83	PROT
	ATOM	1512 CG2 VAL 414 51.169		PROT
	ATOM	1513 C VAL 414 50.630	7.529 44.225 1.00 17.96	PROT
	ATOM	1514 O VAL 414 49.708	8.236 43.829 1.00 30.33	PROT
5	ATOM	1515 N THR 415 51.586	7.969 45.028 1.00 32.51	PROT
-	ATOM	1516 CA THR 415 51.601		PROT
	ATOM		9.529 46.511 1.00 49.75	PROT
	ATOM		3 10.930 46.702 1.00 60.64	PROT
	ATOM	1519 CG2 THR 415 54.03		PROT
10	ATOM		10.387 44.436 1.00 31.44	PROT
10	ATOM		10.251 43.475 1.00 22.01	PROT
	ATOM		11.437 44.607 1.00 24.94	PROT
	ATOM		12.559 43.671 1.00 27.82	PROT
	ATOM		13.164 43.440 1.00 32.98	PROT
15	ATOM		13.747 44.671 1.00 44.74	PROT
	ATOM		13.539 45.251 1.00 44.91	PROT
	ATOM		14.665 45.462 1.00 49.20	PROT
	ATOM		15.000 46.477 1.00 53.14	PROT
	ATOM		14.330 46.373 1.00 41.72	PROT
20	ATOM		12.172 42.328 1.00 29.13	PROT
_,	ATOM		12.660 41.286 1.00 37.24	PROT
	ATOM		11.311 42.350 1.00 18.38	PROT
	ATOM		10.867 41.115 1.00 16.08	PROT
	ATOM		10.029 41.407 1.00 17.89	PROT
25	ATOM	1535 CG PHE 417 47.021	8.997 40.364 1.00 16.15	PROT
	ATOM	1536 CD1 PHE 417 47.980		PROT
	ATOM	1537 CD2 PHE 417 45.800	6 8.971 39.696 1.00 15.49	PROT
	<b>ATOM</b>	1538 CEI PHE 417 47.727	7 7.081 39.087 1.00 19.81	PROT
	ATOM	1539 CE2 PHE 417 45.544		PROT
30	ATOM	1540 CZ PHE 417 46.501	7.064 38.427 1.00 5.25	PROT
	ATOM		11.990 40.187 1.00 14.51	PROT
	ATOM	1542 O PHE 417 48.636	12.119 39.081 1.00 18.44	PROT
	ATOM		12.800 40.640 1.00 21.08	PROT
	ATOM		13.900 39.828 1.00 16.28	PROT
35	ATOM		14.832 40.659 1.00 15.19	PROT
	ATOM		15.746 39.802 1.00 16.60	PROT
	ATOM		5 15.369 38.710 1.00 21.85	PROT
	ATOM		16.557 38.118 1.00 22.53	PROT
	ATOM		14.138 38.170 1.00 16.42	PROT
40	ATOM	1550 CD1 TRP 418 44.999	9 17.107 39.836 1.00 21.01	PROT
	ATOM		5 17.606 38.826 1.00 24.02	PROT
	ATOM	1552 CZ2 TRP 418 42.838	3 16.555 37.010 1.00 24.64	PROT
	ATOM	1553 CZ3 TRP 418 42.925	5 14.135 37.069 1.00 28.80	PROT
	ATOM		7 15.337 36.500 1.00 21.25	PROT
45	ATOM		14.676 39.192 1.00 16.17	PROT
	ATOM		14.764 37.977 1.00 19.51	PROT
	ATOM		15.250 40.007 1.00 19.59	PROT
	ATOM		7 15.274 41.477 1.00 19.81	PROT
	ATOM		7 16.002 39.429 1.00 17.87	PROT
50	ATOM	1560 CB PRO 419 50.720	16.309 40.629 1.00 6.85	PROT

	ATOM	1561 CG PRO 419	49.785 16.326 41.764 1.00 25.11	PROT
	ATOM	1562 C PRO 419	50.578 15.202 38.373 1.00 15.44	PROT
	ATOM	1563 O PRO 419	50.922 15.720 37.315 1.00 24.75	PROT
	ATOM	1564 N LYS 420	50.811 13.932 38.664 1.00 15.10	PROT
5	ATOM	1565 CA LYS 420	51.534 13.056 37.748 1.00 20.59	PROT
_	ATOM	1566 CB LYS 420	51.900 11.746 38.471 1.00 28.85	PROT
	ATOM	1567 CG LYS 420	52,955 11,906 39,577 1,00 30.61	PROT
	ATOM	1568 CD LYS 420	52.907 10.759 40.580 1.00 24.41	PROT
	ATOM	1569 CE LYS 420	54.275 10.493 41.224 1.00 31.94	<b>PROT</b>
10	ATOM	1570 NZ LYS 420	54.485 9.040 41.557 1.00 27.34	PROT
	ATOM	1571 C LYS 420	50.779 12.757 36.445 1.00 17.36	PROT
	ATOM	1572 O LYS 420	51.393 12.439 35.437 1.00 26.28	PROT
	ATOM	1573 N LEU 421	49.455 12.859 36.474 1.00 16.34	PROT
	ATOM	1574 CA LEU 421	48.627 12.614 35.297 1.00 9.38	PROT
15	ATOM	1575 CB LEU 421	47.231 12.139 35.707 1.00 13.22	PROT
13	ATOM	1576 CG LEU 421	46.739 10.818 35.107 1.00 15.75	PROT
	ATOM	1577 CD1 LEU 421	47.919 9.993 34.652 1.00 29.24	PROT
	ATOM	1577 CD1 LEU 421	45.949 10.049 36.135 1.00 12.19	PROT
	ATOM	1579 C LEU 421	48.511 13.866 34.441 1.00 12.61	PROT
20	ATOM	1580 O LEU 421	48.458 13.777 33.223 1.00 17.85	PROT
20	ATOM	1581 N LEU 422	48.451 15.036 35.063 1.00 8.47	PROT
	ATOM	1582 CA LEU 422	48.393 16.254 34.277 1.00 7.21	PROT
	ATOM	1583 CB LEU 422	48.160 17.468 35.164 1.00 2.00	PROT
	ATOM	1584 CG LEU 422	46.941 17.445 36.088 1.00 12.16	PROT
25	ATOM	1585 CD1 LEU 422	47.024 18.660 36.982 1.00 6.96	PROT
23	ATOM	1586 CD2 LEU 422	45.632 17.450 35.313 1.00 2.00	PROT
	ATOM	1587 C LEU 422	49.748 16.365 33.567 1.00 10.59	PROT
	ATOM	1588 O LEU 422	49.851 16.938 32.477 1.00 13.48	PROT
	ATOM	1589 N MET 423	50.786 15.804 34.185 1.00 2.29	PROT
30	ATOM	1590 CA MET 423	52.109 15.821 33.579 1.00 6.50	PROT
50	ATOM	1591 CB MET 423	53.158 15.215 34.514 1.00 2.13	PROT
	ATOM	1592 CG MET 423	53.361 15.968 35.803 1.00 16.33	PROT
	ATOM	1593 SD MET 423	55.075 16.415 36.070 1.00 26.66	PROT
	ATOM	1594 CE MET 423	55.751 14.880 36.623 1.00 20.24	PROT
35	ATOM	1595 C MET 423	52.016 14.966 32.318 1.00 12.20	PROT
55	ATOM	1596 O MET 423	52.741 15.183 31.345 1.00 18.67	PROT
	ATOM	1597 N LYS 424	51.114 13.988 32.352 1.00 7.89	PROT
	ATOM	1598 CA LYS 424	50.907 13.084 31.230 1.00 12.91	PROT
	ATOM	1599 CB LYS 424	49.990 11.924 31.645 1.00 5.14	PROT
40	ATOM	1600 CG LYS 424	50.669 10.579 31.980 1.00 11.76	PROT
40	ATOM	1601 CD LYS 424	52.187 10.590 31.866 1.00 3.70	PROT
	ATOM	1602 CE LYS 424	52.844 10.020 33.113 1.00 7.84	PROT
	ATOM	1603 NZ LYS 424	54.335 9.959 32.995 1.00 25.86	PROT
	ATOM	1604 C LYS 424	50.293 13.840 30.046 1.00 17.44	PROT
15		1605 O LYS 424	50.650 13.596 28.897 1.00 11.72	PROT
45	ATOM	1605 O LYS 424 1606 N VAL 425	49.370 14.756 30.322 1.00 3.16	PROT
	ATOM ATOM	1606 N VAL 425	48.768 15.515 29.249 1.00 2.00	PROT
	ATOM	1607 CA VAL 425 1608 CB VAL 425	47.744 16.532 29.773 1.00 6.77	PROT
	ATOM	1609 CG1 VAL 425	47.744 10.332 29.773 1.00 0.77	PROT
50		1610 CG2 VAL 425	46.381 15.870 29.914 1.00 10.91	PROT
50	ATOM	1010 CG2 VAL 423	TU.JUI 1J.070 47.717 1.00 10.71	1 1/0 1

	ATOM	1611 C VAL 425	49.845 16.274 28.487 1.00 4.83	PROT
	ATOM	1612 O VAL 425	49.853 16.265 27.269 1.00 15.69	PROT
	ATOM	1613 N THR 426	50.753 16.924 29.208 1.00 14.38	PROT
	ATOM	1614 CA THR 426	51.824 17.707 28.593 1.00 12.41	PROT
5	ATOM	1615 CB THR 426	52.713 18.372 29.667 1.00 12.49	PROT
-	ATOM	1616 OG1 THR 426	51.890 19.138 30.552 1.00 11.06	PROT
	ATOM	1617 CG2 THR 426	53.763 19.283 29.015 1.00 2.93	PROT
	ATOM	1618 C THR 426	52.734 16.928 27.653 1.00 15.72	PROT
	ATOM	1619 O THR 426	53.198 17.463 26.651 1.00 14.40	PROT
10	ATOM	1620 N ASP 427	53.000 15.672 27.981 1.00 16.23	PROT
	<b>ATOM</b>	1621 CA ASP 427	53.865 14.843 27.157 1.00 16.35	PROT
	<b>ATOM</b>	1622 CB ASP 427	54.342 13.630 27.950 1.00 19.48	PROT
	ATOM	1623 CG ASP 427	55.337 13.997 29.029 1.00 18.96	PROT
	ATOM	1624 OD1 ASP 427	55.874 15.125 29.010 1.00 8.75	PROT
15	ATOM	1625 OD2 ASP 427	55.579 13.145 29.902 1.00 24.25	PROT
	ATOM	1626 C ASP 427	53.155 14.381 25.891 1.00 20.52	PROT
	ATOM	1627 O ASP 427	53.793 14.164 24.856 1.00 25.69	PROT
	<b>ATOM</b>	1628 N LEU 428	51.838 14.218 25.986 1.00 5.49	PROT
	ATOM	1629 CA LEU 428	51.040 13.815 24.849 1.00 2.00	PROT
20	<b>ATOM</b>	1630 CB LEU 428	49.634 13.470 25.301 1.00 2.00	PROT
	<b>ATOM</b>	1631 CG LEU 428	49.579 12.127 26.028 1.00 2.00	PROT
	<b>ATOM</b>	1632 CD1 LEU 428	48.184 11.789 26.481 1.00 2.00	PROT
	<b>ATOM</b>	1633 CD2 LEU 428	50.088 11.080 25.108 1.00 2.00	PROT
	<b>ATOM</b>	1634 C LEU 428	51.019 14.987 23.881 1.00 7.72	PROT
25	<b>ATOM</b>	1635 O LEU 428	51.072 14.800 22.666 1.00 9.22	PROT
	ATOM	1636 N ARG 429	50.961 16.197 24.432 1.00 10.07	PROT
	ATOM	1637 CA ARG 429	50.948 17.438 23.659 1.00 7.97	PROT
	ATOM	1638 CB ARG 429	50.799 18.642 24.583 1.00 18.55	PROT
	ATOM	1639 CG ARG 429	49.548 18.634 25.429 1.00 14.80	PROT
30	ATOM	1640 CD ARG 429	48.588 19.674 24.935 1.00 32.08	PROT
	ATOM	1641 NE ARG 429	47.508 19.923 25.880 1.00 42.46	PROT
	ATOM	1642 CZ ARG 429	46.226 19.673 25.631 1.00 48.51	PROT
	ATOM	1643 NH1 ARG 429		PROT
	ATOM	1644 NH2 ARG 429		PROT
35	ATOM	1645 C ARG 429	52.260 17.557 22.919 1.00 11.77	PROT
	ATOM	1646 O ARG 429	52.298 17.904 21.737 1.00 28.66	PROT
	ATOM	1647 N MET 430	53.343 17.270 23.629 1.00 20.26	PROT
	ATOM	1648 CA MET 430	54.671 17.328 23.042 1.00 21.06	PROT
	ATOM	1649 CB MET 430	55.738 17.015 24.100 1.00 30.24	PROT
40	ATOM	1650 CG MET 430	56.061 18.165 25.056 1.00 34.66	PROT
	ATOM	1651 SD MET 430	55.727 19.795 24.373 1.00 35.91	PROT
	ATOM	1652 CE MET 430	56.839 19.814 22.978 1.00 32.52	PROT
	ATOM	1653 C MET 430	54.735 16.302 21.925 1.00 18.70	PROT
	ATOM	1654 O MET 430	55.287 16.560 20.860 1.00 16.59	PROT
45	ATOM	1655 N ILE 431	54.161 15.133 22.182 1.00 15.38	PROT
	ATOM	1656 CA ILE 431	54.144 14.069 21.196 1.00 15.85	PROT
	ATOM	1657 CB ILE 431	53.326 12.859 21.705 1.00 13.76	PROT
	ATOM	1658 CG2 ILE 431	52.727 12.084 20.539 1.00 11.11	PROT
	ATOM	1659 CG1 ILE 431	54.239 11.924 22.489 1.00 11.72	PROT
50	ATOM	1660 CD1 ILE 431	53.552 11.224 23.615 1.00 16.22	PROT

	ATOM	1661 C ILE 431 53.538 14.609 19.904 1.00 18.49 PROT	
	ATOM	1662 O ILE 431 54.134 14.483 18.839 1.00 17.36 PROT	
	ATOM	1663 N GLY 432 52.361 15.220 20.003 1.00 2.00 PROT	
	ATOM	1664 CA GLY 432 51.721 15.772 18.831 1.00 2.00 PROT	
5	ATOM	1665 C GLY 432 52.542 16.851 18.148 1.00 10.55 PROT	
-	ATOM	1666 O GLY 432 52.707 16.834 16.936 1.00 9.60 PROT	
	ATOM	1667 N ALA 433 53.043 17.805 18.926 1.00 11.17 PROT	
	ATOM	1668 CA ALA 433 53.855 18.884 18.385 1.00 2.00 PROT	
	ATOM	1669 CB ALA 433 54.326 19.771 19.506 1.00 2.00 PROT	
10	ATOM	1670 C ALA 433 55.050 18.285 17.646 1.00 6.43 PROT	
	ATOM	1671 O ALA 433 55.493 18.789 16.623 1.00 11.71 PROT	
	ATOM	1672 N CYS 434 55.579 17.197 18.179 1.00 15.71 PROT	
	ATOM	1673 CA CYS 434 56.715 16.534 17.573 1.00 13.44 PROT	
	ATOM	1674 CB CYS 434 57.228 15.464 18.518 1.00 14.76 PROT	
15	ATOM	1675 SG CYS 434 58.910 15.703 18.985 1.00 20.82 PROT	
	ATOM	1676 C CYS 434 56.269 15.902 16.264 1.00 9.28 PROT	
	ATOM	1677 O CYS 434 56.969 15.948 15.256 1.00 8.50 PROT	
	ATOM	1678 N HIS 435 55.091 15.300 16.298 1.00 11.04 PROT	
	ATOM	1679 CA HIS 435 54.533 14.657 15.122 1.00 11.30 PROT	
20	ATOM	1680 CB HIS 435 53.142 14.132 15.438 1.00 4.30 PROT	
	ATOM	1681 CG HIS 435 52.480 13.460 14.283 1.00 13.68 PROT	
	ATOM	1682 CD2 HIS 435 52.751 12.288 13.662 1.00 4.72 PROT	
	<b>ATOM</b>	1683 ND1 HIS 435 51.358 13.976 13.666 1.00 5.53 PROT	
	<b>ATOM</b>	1684 CE1 HIS 435 50.966 13.147 12.717 1.00 12.84 PROT	
25	<b>ATOM</b>	1685 NE2 HIS 435 51.794 12.116 12.694 1.00 15.77 PROT	
	<b>ATOM</b>	1686 C HIS 435 54.482 15.661 13.973 1.00 8.50 PROT	
	<b>ATOM</b>	1687 O HIS 435 54.941 15.370 12.869 1.00 14.82 PROT	
	ATOM	1688 N ALA 436 53.938 16.844 14.245 1.00 5.74 PROT	
	<b>ATOM</b>	1689 CA ALA 436 53.843 17.905 13.252 1.00 2.00 PROT	
30	ATOM	1690 CB ALA 436 53.632 19.241 13.942 1.00 2.00 PROT	
	ATOM	1691 C ALA 436 55.121 17.934 12.406 1.00 8.68 PROT	
	ATOM	1692 O ALA 436 55.080 17.712 11.193 1.00 15.14 PROT	
	ATOM	1693 N SER 437 56.256 18.189 13.047 1.00 6.82 PROT	
	ATOM	1694 CA SER 437 57.522 18.226 12.337 1.00 9.05 PROT	
35	<b>ATOM</b>	1695 CB SER 437 58.671 18.511 13.295 1.00 2.00 PROT	
	ATOM	1696 OG SER 437 59.593 19.406 12.699 1.00 21.18 PROT	
	ATOM	1697 C SER 437 57.758 16.896 11.637 1.00 15.18 PROT	
	ATOM	1698 O SER 437 58.076 16.849 10.445 1.00 19.33 PROT	
	ATOM	1699 N ARG 438 57.607 15.805 12.373 1.00 16.98 PROT	
40	ATOM	1700 CA ARG 438 57.799 14.501 11.766 1.00 16.98 PROT	
	ATOM	1701 CB ARG 438 57.294 13.409 12.702 1.00 24.77 PROT	
	ATOM	1702 CG ARG 438 58.006 12.086 12.534 1.00 33.76 PROT	
	ATOM		
	ATOM		
45	ATOM	1705 CZ ARG 438 61.505 11.504 11.423 1.00 25.21 PROT	
	ATOM	1706 NH1 ARG 438 62.077 10.641 10.603 1.00 39.58 PROT	
	ATOM		
	ATOM		
50	ATOM ATOM	1709 O ARG 438 57.563 14.008 9.424 1.00 15.57 PROT 1710 N PHE 439 55.781 14.893 10.484 1.00 16.75 PROT	

	ATOM	1711 CA PHE 439	54.933 14.878 9.303 1.00 21.63	PROT
	ATOM	1712 CB PHE 439	53.603 15.575 9.574 1.00 17.84	PROT
	ATOM	1713 CG PHE 439	52.597 15.364 8.490 1.00 20.60	PROT
	ATOM	1714 CD1 PHE 439	52.042 14.103 8.279 1.00 30.60	PROT
5	ATOM	1715 CD2 PHE 439	52.265 16.394 7.622 1.00 14.95	PROT
	ATOM	1716 CE1 PHE 439	51.175 13.867 7.206 1.00 29.12	PROT
	ATOM	1717 CE2 PHE 439	51.404 16.173 6.552 1.00 25.18	PROT
	ATOM	1718 CZ PHE 439	50.860 14.905 6.341 1.00 27.82	PROT
	ATOM	1719 C PHE 439	55.620 15.548 8.130 1.00 28.17	PROT
10	ATOM	1720 O PHE 439	55.512 15.095 6.987 1.00 28.83	PROT
	ATOM	1721 N LEU 440	56.328 16.633 8.427 1.00 26.77	PROT
	ATOM	1722 CA LEU 440	57.055 17.382 7.418 1.00 24.66	PROT
	ATOM	1723 CB LEU 440	57.555 18.696 8.005 1.00 10.80	PROT
	ATOM	1724 CG LEU 440	56.501 19.658 8.541 1.00 8.60	PROT
15	ATOM	1725 CD1 LEU 440	57.152 20.985 8.855 1.00 17.69	PROT
	ATOM	1726 CD2 LEU 440	55.410 19.847 7.522 1.00 15.71	PROT
	ATOM	1727 C LEU 440	58.245 16.578 6.912 1.00 29.61	PROT
	ATOM	1728 O LEU 440	58.506 16.526 5.718 1.00 32.37	PROT
	ATOM		58.971 15.954 7.830 1.00 28.12	PROT
20	ATOM	1730 CA HIS 441	60.140 15.172 7.460 1.00 28.51	PROT
	ATOM	1731 CB HIS 441	60.783 14.564 8.705 1.00 36.77	PROT
	ATOM		59.724 14.081 6.497 1.00 31.94	PROT
	ATOM	1733 O HIS 441	60.461 13.725 5.579 1.00 49.29	PROT
	ATOM	1734 N MET 442	58.533 13.545 6.711 1.00 41.16	PROT
25	ATOM	1735 CA MET 442	58.033 12.487 5.854 1.00 39.99	PROT
	ATOM	1736 CB MET 442	56.871 11.776 6.551 1.00 38.32	PROT
	ATOM	1737 CG MET 442	57.263 11.122 7.860 1.00 19.20	PROT
	ATOM	1738 SD MET 442	55.859 10.350 8.675 1.00 38.06	PROT
	ATOM	1739 CE MET 442	54.906 11.767 9.073 1.00 21.45	PROT
30	ATOM	1740 C MET 442	57.599 13.031 4.495 1.00 35.68	PROT
	ATOM	1741 O MET 442	57.887 12.431 3.461 1.00 27.43	PROT
	<b>ATOM</b>	1742 N LYS 443	56.920 14.175 4.503 1.00 34.17	PROT
	<b>ATOM</b>	1743 CA LYS 443	56.447 14.796 3.268 1.00 34.33	PROT
	<b>ATOM</b>	1744 CB LYS 443	55.767 16.129 3.574 1.00 21.68	PROT
35	<b>ATOM</b>	1745 CG LYS 443	54.303 15.989 3.953 1.00 26.95	PROT
	<b>ATOM</b>	1746 CD LYS 443	53.497 17.231 3.602 1.00 30.78	PROT
	<b>ATOM</b>	1747 CE LYS 443	52.204 16.848 2.861 1.00 56.06	PROT
	<b>ATOM</b>	1748 NZ LYS 443	50.931 17.261 3.564 1.00 45.26	PROT
	ATOM	1749 C LYS 443	57.570 15.007 2.251 1.00 37.81	PROT
40	ATOM	1750 O LYS 443	57.325 15.049 1.041 1.00 38.26	PROT
	ATOM	1751 N VAL 444	58.798 15.130 2.741 1.00 25.12	PROT
	<b>ATOM</b>	1752 CA VAL 444	59.942 15.318 1.867 1.00 25.43	PROT
	<b>ATOM</b>	1753 CB VAL 444	60.802 16.531 2.334 1.00 29.15	PROT
	<b>ATOM</b>	1754 CG1 VAL 444	59.893 17.621 2.861 1.00 29.48	PROT
45	ATOM	1755 CG2 VAL 444	61.785 16.121 3.419 1.00 36.65	PROT
	<b>ATOM</b>	1756 C VAL 444	60.786 14.042 1.825 1.00 30.03	PROT
	<b>ATOM</b>	1757 O VAL 444	62.009 14.099 1.698 1.00 39.43	PROT
	<b>ATOM</b>	1758 N GLU 445	60.127 12.888 1.903 1.00 39.84	PROT
	ATOM	1759 CA GLU 445	60.842 11.612 1.896 1.00 43.07	PROT
50	ATOM	1760 CB GLU 445	61.429 11.360 3.282 1.00 50.55	PROT

	ATOM	1761 CG GLU 445	62.399 10.203 3.351 1.00 77.00	PROT
	ATOM	1762 CD GLU 445	63.569 10.495 4.267 1.00 98.21	PROT
	ATOM	1763 OE1 GLU 445	64.251 9.538 4.701 1.00100.00	PROT
	ATOM	1764 OE2 GLU 445	63.804 11.690 4.554 1.00100.00	
5	ATOM	1765 C GLU 445	59.989 10.408 1.491 1.00 43.41	PROT
	ATOM	1766 O GLU 445	60.466 9.274 1.511 1.00 48.80	PROT
	ATOM	1767 N CYS 446	58.731 10.644 1.137 1.00 38.17	PROT
	ATOM	1768 CA CYS 446	57.852 9.548 0.743 1.00 41.38	PROT
	ATOM	1769 CB CYS 446	57.066 9.035 1.965 1.00 40.61	PROT
10	ATOM	1770 SG CYS 446	58.062 8.276 3.320 1.00 44.73	PROT
	ATOM	1771 C CYS 446	56.886 10.003 -0.362 1.00 45.83	PROT
	ATOM	1772 O CYS 446	56.466 11.184 -0.323 1.00 44.17	PROT
	ATOM	1773 OT CYS 446	56.570 9.180 -1.259 1.00 40.79	PROT
	ATOM	1774 CB GLU 449	52.635 12.140 -2.649 1.00 28.60	PROT
15	ATOM	1775 C GLU 449	52.019 10.014 -1.526 1.00 38.06	PROT
	ATOM	1776 O GLU 449	50.873 10.220 -1.935 1.00 43.52	PROT
	ATOM	1777 N GLU 449	54.378 10.460 -2.167 1.00 17.78	PROT
	ATOM	1778 CA GLU 449	53.105 11.069 -1.689 1.00 33.80	PROT
	ATOM	1779 N LEU 450	52.387 8.880 -0.936 1.00 46.88	PROT
20	ATOM	1780 CA LEU 450	51.432 7.808 -0.696 1.00 52.62	PROT
	ATOM	1781 CB LEU 450	52.101 6.436 -0.850 1.00 57.50	PROT
	ATOM	1782 CG LEU 450	53.338 6.066 -0.028 1.00 59.81	PROT
	ATOM	1783 CD1 LEU 450	53.613 4.573 -0.198 1.00 51.33	PROT
	ATOM	1784 CD2 LEU 450	54.544 6.890 -0.473 1.00 57.03	PROT
25	ATOM	1785 C LEU 450	50.850 7.970 0.711 1.00 50.65	PROT
	ATOM	1786 O LEU 450	50.965 7.091 1.569 1.00 38.49	PROT
	ATOM	1787 N PHE 451	50.225 9.123 0.923 1.00 32.24	PROT
	ATOM	1788 CA PHE 451	49.602 9.478 2.188 1.00 32.64	PROT
	ATOM	1789 CB PHE 451	50.091 10.857 2.648 1.00 56.06	PROT
30	ATOM	1790 CG PHE 451	51.534 10.895 3.056 1.00 61.73	PROT
	ATOM	1791 CD1 PHE 451	52.523 10.366 2.235 1.00 66.92	PROT
	ATOM	1792 CD2 PHE 451	51.905 11.486 4.256 1.00 58.76	PROT
	ATOM	1793 CE1 PHE 451	53.860 10.430 2.604 1.00 69.17	PROT
	ATOM	1794 CE2 PHE 451	53.231 11.556 4.635 1.00 61.48	PROT
35	ATOM	1795 CZ PHE 451	54.214 11.028 3.809 1.00 71.95	PROT
	ATOM	1796 C PHE 451	48.081 9.548 2.025 1.00 30.67	PROT
	ATOM	1797 O PHE 451	47.571 10.429 1.324 1.00 38.49	PROT
	ATOM	1798 N PRO 452	47.336 8.627 2.672 1.00 19.14	PROT
40	ATOM	1799 CD PRO 452	47.774 7.495 3.510 1.00 24.21	PROT
40	ATOM	1800 CA PRO 452	45.881 8.672 2.538 1.00 5.88	PROT
	ATOM	1801 CB PRO 452	45.397 7.742 3.633 1.00 16.92	PROT
	ATOM	1802 CG PRO 452	46.496 6.737 3.761 1.00 16.91	PROT
	ATOM	1803 C PRO 452	45.354 10.090 2.687 1.00 15.15	PROT
15	ATOM	1804 O PRO 452	45.879 10.886 3.463 1.00 22.59	PROT
45	ATOM	1805 N PRO 453	44.315 10.429 1.920 1.00 18.37	PROT
	ATOM	1806 CD PRO 453	43.653 9.540 0.951 1.00 3.83	PROT
	ATOM	1807 CA PRO 453	43.710 11.766 1.960 1.00 14.00	PROT PROT
	ATOM	1808 CB PRO 453	42.502 11.649 1.032 1.00 20.04 42.316 10.163 0.807 1.00 19.43	PROT
50	ATOM ATOM	1809 CG PRO 453 1810 C PRO 453	43.321 12.277 3.346 1.00 14.70	PROT
50	VI OM	1010 C 1 VO 400	12.221 12.277 2.270 1.00 14.70	INOI

	ATOM	1811 O PRO 453	43.609 13.422 3.682 1.00 9.70	PROT
	<b>ATOM</b>	1812 N LEU 454	42.667 11.446 4.152 1.00 25.39	PROT
	<b>ATOM</b>	1813 CA LEU 454	42.261 11.886 5.491 1.00 28.61	PROT
	<b>ATOM</b>	1814 CB LEU 454	41.463 10.804 6.217 1.00 17.29	PROT
5	ATOM	1815 CG LEU 454	40.893 11.224 7.572 1.00 9.05	PROT
	ATOM	1816 CD1 LEU 454	40.174 12.547 7.435 1.00 17.23	PROT
	ATOM	1817 CD2 LEU 454	39.946 10.148 8.079 1.00 8.05	PROT
	ATOM	1818 C LEU 454	43,479 12,234 6.316 1.00 23.36	PROT
	ATOM	1819 O LEU 454	43.484 13.225 7.037 1.00 10.99	PROT
10	ATOM	1820 N PHE 455	44.503 11.394 6.205 1.00 14.26	PROT
	ATOM	1821 CA PHE 455	45.769 11.595 6.902 1.00 15.33	PROT
	ATOM	1822 CB PHE 455	46.761 10.496 6.501 1.00 26.32	PROT
	ATOM	1823 CG PHE 455	48.138 10.644 7.108 1.00 43.03	PROT
	ATOM	1824 CD1 PHE 455	48.305 11.094 8.414 1.00 43.52	PROT
15	ATOM	1825 CD2 PHE 455	49.270 10.282 6.380 1.00 41.44	PROT
13	ATOM	1826 CE1 PHE 455	49.576 11.176 8.987 1.00 37.77	PROT
	ATOM	1827 CE2 PHE 455	50.536 10.363 6.947 1.00 49.43	PROT
	ATOM	1828 CZ PHE 455	50.686 10.811 8.255 1.00 39.99	PROT
	ATOM	1829 C PHE 455	46.313 12.956 6.500 1.00 19.37	PROT
20	ATOM	1830 O PHE 455	46.945 13.646 7.298 1.00 29.31	PROT
20	ATOM	1831 N LEU 456	46.048 13.345 5.257 1.00 17.16	PROT
	ATOM	1832 CA LEU 456	46.527 14.625 4.750 1.00 20.15	PROT
	ATOM	1833 CB LEU 456	46.572 14.603 3.218 1.00 35.14	PROT
	ATOM	1834 CG LEU 456	47.593 13.660 2.568 1.00 40.45	PROT
25	ATOM	1835 CD1 LEU 456	47.233 13.456 1.116 1.00 44.38	PROT
	ATOM	1836 CD2 LEU 456	48.990 14.234 2.680 1.00 34.88	PROT
	ATOM	1837 C LEU 456	45.680 15.800 5.226 1.00 20.37	PROT
	ATOM	1838 O LEU 456	46.207 16.866 5.548 1.00 29.61	PROT
	ATOM	1839 N GLU 457	44.367 15.607 5.280 1.00 13.06	PROT
30	ATOM	1840 CA GLU 457	43.483 16.675 5.713 1.00 14.14	PROT
	ATOM	1841 CB GLU 457	42.037 16.256 5.516 1.00 29.57	PROT
	ATOM	1842 C GLU 457	43.731 17.058 7.173 1.00 14.95	PROT
	ATOM	1843 O GLU 457	43.771 18.237 7.514 1.00 15.98	PROT
	ATOM	1844 N VAL 458	43.901 16.051 8.026 1.00 26.34	PROT
35	ATOM	1845 CA VAL 458	44.143 16.260 9.455 1.00 24.39	PROT
	ATOM	1846 CB VAL 458	44.219 14.910 10.208 1.00 20.14	PROT
	ATOM	1847 CG1 VAL 458	44.882 15.102 11.554 1.00 22.01	PROT
	ATOM	1848 CG2 VAL 458	42.831 14.341 10.400 1.00 28.11	PROT
	ATOM	1849 C VAL 458	45.417 17.039 9.778 1.00 21.50	PROT
40	ATOM	1850 O VAL 458	45.364 18.062 10.439 1.00 18.85	PROT
	ATOM	1851 N PHE 459	46.557 16.546 9.308 1.00 16.05	PROT
	ATOM	1852 CA PHE 459	47.840 17.174 9.586 1.00 20.28	PROT
	ATOM	1853 CB PHE 459	48.862 16.072 9.846 1.00 20.26	PROT
	ATOM	1854 CG PHE 459	48.389 15.055 10.833 1.00 27.22	PROT
45	ATOM	1855 CD1 PHE 459	47.917 13.822 10.408 1.00 28.01	PROT
	ATOM	1856 CD2 PHE 459	48.390 15.339 12.204 1.00 40.66	PROT
	ATOM	1857 CE1 PHE 459	47.447 12.876 11.334 1.00 21.78	PROT
	ATOM	1858 CE2 PHE 459	47.922 14.402 13.140 1.00 25.98	PROT
	ATOM	1859 CZ PHE 459	47.450 13.172 12.702 1.00 17.63	PROT
50	ATOM	1860 C PHE 459	48.381 18.152 8.540 1.00 23.03	PROT

	ATOM	1861 O PHE 459 49.601 18.311 8.416 1.00 27.	34 PROT
	ATOM	1862 N GLU 460 47.480 18.816 7.815 1.00 33.	88 PROT
	ATOM	1863 CA GLU 460 47.846 19.774 6.767 1.00 36	6.60 PROT
	ATOM	1864 CB GLU 460 48.930 20.732 7.257 1.00 46	5.04 PROT
5	ATOM	1865 CG GLU 460 48.406 21.899 8.054 1.00 67	7.27 PROT
	ATOM	1866 CD GLU 460 47.298 22.636 7.339 1.00 71	1.34 PROT
	ATOM	1867 OE1 GLU 460 47.448 23.859 7.121 1.00 7	1.99 PROT
	ATOM	1868 OE2 GLU 460 46.280 21.993 6.998 1.00 7	2.73 PROT
	ATOM	1869 C GLU 460 48.353 19.037 5.535 1.00 46.	
10	ATOM	1870 O GLU 460 48.642 17.829 5.655 1.00 51.	
	<b>ATOM</b>	1871 OT GLU 460 48.461 19.669 4.462 1.00 60	
	<b>ATOM</b>	1872 C1 GC1 1 47.011 4.539 15.912 1.00 29.3	
	<b>ATOM</b>	1873 C2 GC1 1 51.292 6.537 13.571 1.00 17.1	1 LIGA
	<b>ATOM</b>	1874 C3 GC1 1 47.393 4.205 14.573 1.00 33.7	
15	<b>ATOM</b>	1875 C4 GC1 1 52.119 6.409 12.400 1.00 19.7	
	<b>ATOM</b>	1876 C5 GC1 1 48.689 4.481 14.089 1.00 25.0	
	<b>ATOM</b>	1877 C6 GC1 1 52.344 7.525 11.539 1.00 17.5	
	ATOM	1878 C7 GC1 1 49.684 5.122 14.949 1.00 23.9	
	<b>ATOM</b>	1879 C8 GC1 1 51.722 8.778 11.873 1.00 20.2	
20	ATOM	1880 C9 GC1 1 49.283 5.452 16.318 1.00 18.1	
	ATOM	1881 C10 GC1 1 50.906 8.928 13.018 1.00 15.	
	<b>ATOM</b>	1882 C11 GC1 1 47.973 5.163 16.779 1.00 30.	
	ATOM	1883 C12 GC1 1 50.696 7.827 13.850 1.00 25.	
	ATOM	1884 O5 GC1 1 45.700 4.254 16.325 1.00 28.6	
25	ATOM	1885 C14 GC1 1 53.198 7.459 10.291 1.00 20.	
	ATOM	1886 C15 GC1 1 45.305 3.866 17.666 1.00 18.	
	ATOM	1887 C16 GC1 1 52.423 6.824 9.131 1.00 17.2	
	ATOM	1888 C17 GC1 1 43.816 4.078 17.872 1.00 21.	
	ATOM	1889 C18 GC1 1 54.514 6.689 10.543 1.00 24.	
30	ATOM	1890 C19 GC1 1 48.994 4.093 12.664 1.00 33.	
	ATOM	1891 C20 GC1 1 50.243 6.110 17.278 1.00 27.	
	ATOM	1892 O1 GC1 1 51.902 9.861 11.086 1.00 23.3	
	ATOM	1893 C21 GC1 1 51.026 5.430 14.458 1.00 22.	
	ATOM	1894 O3 GC1 1 43.147 3.117 18.247 1.00 18.0	
35	ATOM	1895 O4 GC1 1 43.331 5.204 17.665 1.00 28.2	27 LIGA
	END		

WO 99/26966

## **APPENDIX 8**

## TRBGC1.PDB

REMARK TR-beta GC-2 Full length numbering

REMARK refinement resolution: 100.00 - 2.40 A starting r= 0.2602 free\_r= 0.2960

REMARK final r=0.2532 free\_r=0.2894

REMARK sg= P3(1)21 a= 68.9 b= 68.9 c= 131.5 alpha= 90 beta= 90 gamma= 120

REMARK theoretical total number of refl. in resol. range: 14710 (100.0 %)

REMARK number of unobserved reflections (no entry or |F|=0): 336 (2.3 %)

REMARK number of reflections rejected:

0(0.0%)

10 REMARK total number of reflections used:

14374 ( 97.7 % )

REMARK number of reflections in working set:

13656 (92.8%)

PCT/US98/25296

REMARK number of reflections in test set:

718 ( 4.9 % )

REMARK

REMARK ALA 199 to ALA 201 from His-tag

15 REMARK

REMARK Four cacodylate-modified cysteines (CYA)

REMARK Cys294, Cys298, Cys388, Cys434

REMARK cacodylate modeled as single arsenic atom

REMARK

20 REMARK side chain of certain residues modeled as ALA due to poor density;

REMARK however, residue name reflects true residue for clarity

REMARK

REMARK amino acid sequence confirmed,

REMARK differing from that reported by Weinberger et. al.

25 REMARK in the following codons:

REMARK 243 Pro - Arg

REMARK 337 lle - Thr

REMARK 451 Leu - Phe

REMARK as reported by Sakurai et. al.

30 REMARK note also correction of initiation codon,

REMARK yielding a polypeptide of 461 amino acids

JRNL AUTH A.SAKURAI, A.NAKAI, L.J. DEGROOT

JRNL TITL STRUCTURAL ANALYSIS OF HUMAN THYROID HORMONE RECEPTOR

35 JRNL TITL2 BETA GENE

JRNL REF MOL.CELL.ENDO. V.71 1990

JRNL AUTH

C.WEINBERGER,

C.C.THOMPSON,R.LEBO,D.J.GRUOL,R.M.EVANS

JRNL TITL THE C-ERB-A GENE ENCODES A THYROID HORMONE

40 RECEPTOR

JRNL REF NATURE

V.324 6098 1986

ATOM 1 CB ALA 199 36.564 26.104 43.169 1.00 73.87

ATOM 2 C ALA 199 34.723 26.996 44.613 1.00 78.22

ATOM 3 O ALA 199 34.741 28.230 44.568 1.00 81.84

45 ATOM 4 N ALA 199 34.389 26.744 42.166 1.00 77.76

ATOM 5 CA ALA 199 35.048 26.165 43.375 1.00 77.98

ATOM 6 N ALA 200 34.428 26.309 45.713 1.00 77.78

```
34.098 26.961 46.984 1.00 77.03
              7 CA ALA 200
    ATOM
                               32.761 27.693 46.865 1.00 79.04
              8 CB ALA 200
    ATOM
                              34.028 25.897 48.084 1.00 75.79
              9 C ALA 200
    ATOM
                              34.877 25.857 48.978 1.00 71.58
             10 O ALA 200
    ATOM
                               33.005 25.050 48.010 1.00 73.70
             11 N ALA 201
    ATOM
                               32.838 23.968 48.972 1.00 70.15
             12 CA ALA 201
    ATOM
                               31.468 23.328 48.809 1.00 71.16
             13 CB ALA 201
    ATOM
                              33.934 22.963 48.642 1.00 67.54
             14 C ALA 201
    ATOM
             15 O ALA 201
                               34.218 22.044 49.413 1.00 67.14
    ATOM
             16 N GLU 202
                               34.540 23.164 47.476 1.00 62.05
10
    ATOM
                               35.624 22.325 46.975 1.00 59.45
             17 CA GLU 202
    ATOM
                               35.835 22.621 45.482 1.00 55.12
             18 CB GLU 202
    ATOM
                                36.820 21.716 44.749 1.00 56.25
    ATOM
             19 CG GLU 202
                                36.382 20.260 44.723 1.00 54.99
             20 CD GLU 202
    ATOM
             21 OE1 GLU 202
                                35.216 19.990 44.361 1.00 53.83
     ATOM
15
                                37.210 19.385 45.050 1.00 59.90
             22 OE2 GLU 202
     ATOM
                               36.885 22.674 47.770 1.00 55.96
             23 C GLU 202
    ATOM
                               37.472 21.823 48.435 1.00 52.90
    ATOM
             24 O GLU 202
                               37.282 23.943 47.698 1.00 54.95
     ATOM
             25 N GLU 203
                               38.464 24.434 48.390 1.00 55.59
20
     ATOM
             26 CA GLU 203
             27 CB GLU 203
                               38.632 25.924 48.126 1.00 53.21
     ATOM
             28 C GLU 203
                               38.415 24.171 49.894 1.00 56.30
     ATOM
             29 O GLU 203
                               39.445 23.948 50.526 1.00 58.70
     ATOM
     ATOM
             30 N LEU 204
                               37.213 24.193 50.462 1.00 57.14
             31 CA LEU 204
                               37.038 23.966 51.893 1.00 56.93
     ATOM
25
                               35.658 24.465 52.338 1.00 58.31
             32 CB LEU 204
     ATOM
                               35.348 24.508 53.839 1.00 51.69
             33 CG LEU 204
     ATOM
                                36.314 25.446 54.549 1.00 44.38
             34 CD1 LEU 204
     ATOM
             35 CD2 LEU 204
                                33.920 24.986 54.039 1.00 52.44
     ATOM
                               37.198 22.489 52.246 1.00 58.20
     ATOM
             36 C LEU 204
30
     ATOM
             37 O LEU 204
                               37.831 22.155 53.252 1.00 58.99
                               36.620 21.607 51.431 1.00 58.26
             38 N GLN 205
     ATOM
                                36.736 20.167 51.657 1.00 55.38
             39 CA GLN 205
     ATOM
             40 CB GLN 205
                                35.993 19.377 50.584 1.00 54.52
     ATOM
                                34.498 19.324 50.741 1.00 53.33
             41 CG GLN 205
35
     ATOM
             42 CD GLN 205
                                33.854 18.520 49.629 1.00 53.40
     ATOM
             43 OE1 GLN 205
                                33.850 18.939 48.473 1.00 51.68
     ATOM
                                33.325 17.352 49.968 1.00 51.34
             44 NE2 GLN 205
     ATOM
                               38.200 19.775 51.608 1.00 55.05
             45 C GLN 205
     ATOM
                               38.665 18.964 52.407 1.00 53.63
     ATOM
             46 O GLN 205
40
             47 N LYS 206
                               38.918 20.348 50.648 1.00 53.55
     ATOM
             48 CA LYS 206
                               40.337 20.078 50.493 1.00 57.40
     ATOM
                               40.896 20.814 49.269 1.00 58.94
             49 CB LYS 206
     ATOM
              50 CG LYS 206
                                40.300 20.375 47.941 1.00 67.73
     ATOM
                                40.921 21.141 46.781 1.00 72.50
              51 CD LYS 206
45
     ATOM
                               40.346 20.695 45.445 1.00 75.60
     ATOM
              52 CE LYS 206
                               40.945 21.445 44.304 1.00 77.08
              53 NZ LYS 206
     ATOM
                               41.053 20.559 51.747 1.00 53.98
              54 C LYS 206
     ATOM
                               41.905 19.866 52.300 1.00 53.49
              55 O LYS 206
     ATOM
                               40.680 21.757 52.184 1.00 53.61
             56 N SER 207
50
     ATOM
```

	ATOM	57 CA SER 207 41.254 22.386 53.364 1.00 5	1.49
	ATOM	58 CB SER 207 40.546 23.715 53.619 1.00 5	1.01
	ATOM	59 OG SER 207 41.108 24.383 54.731 1.00 6	53.00
	ATOM	60 C SER 207 41.178 21.502 54.616 1.00 49	).49
5	ATOM	61 O SER 207 42.073 21.538 55.465 1.00 4°	
-	ATOM	62 N ILE 208 40.117 20.707 54.725 1.00 44	.39
	ATOM	63 CA ILE 208 39.938 19.829 55.874 1.00 4	
	ATOM	64 CB ILE 208 38.421 19.627 56.174 1.00 44	
	ATOM	65 CG2 ILE 208 38.226 18.801 57.445 1.00 4	
10	ATOM	66 CG1 ILE 208 37.766 20.993 56.385 1.00 4	
	ATOM	67 CD1 ILE 208 36.266 20.941 56.567 1.00 4	
	ATOM	68 C ILE 208 40.614 18.477 55.643 1.00 47	
	ATOM	69 O ILE 208 40.735 17.666 56.562 1.00 49	
	ATOM	70 N GLY 209 41.059 18.238 54.412 1.00 5	
15	ATOM	71 CA GLY 209 41.728 16.983 54.107 1.00	
15	ATOM	72 C GLY 209 40.813 15.896 53.573 1.00 4	
	ATOM	73 O GLY 209 41.203 14.730 53.485 1.00 4	
	ATOM	74 N HIS 210 39.582 16.274 53.237 1.00 46	
	ATOM	75 CA HIS 210 38.622 15.326 52.686 1.00 4	
20	ATOM	76 CB HIS 210 37.200 15.739 53.068 1.00 4	
	ATOM	77 C HIS 210 38.796 15.350 51.162 1.00 45	
	ATOM	78 O HIS 210 38.924 16.420 50.566 1.00 41	
	ATOM	79 N LYS 211 38.829 14.176 50.545 1.00 4	
	ATOM	80 CA LYS 211 38.991 14.095 49.090 1.00 4	
25	ATOM	81 CB LYS 211 39.892 12.910 48.715 1.00 4	6.72
	ATOM	82 CG LYS 211 41.210 12.815 49.497 1.00 5	6.48
	ATOM	83 CD LYS 211 42.068 14.089 49.486 1.00 6	50.93
	ATOM	84 CE LYS 211 42.562 14.496 48.103 1.00 6	1.95
	ATOM	85 NZ LYS 211 41.485 15.024 47.218 1.00 6	9.93
30	ATOM	86 C LYS 211 37.609 13.917 48.473 1.00 35	5.68
	ATOM	87 O LYS 211 37.019 12.847 48.557 1.00 33	3.58
	ATOM	88 N PRO 212 37.077 14.972 47.828 1.00 3	5.64
	<b>ATOM</b>	89 CD PRO 212 37.654 16.304 47.584 1.00 C	38.60
	ATOM	90 CA PRO 212 35.748 14.896 47.211 1.00 C	38.35
35	ATOM	91 CB PRO 212 35.537 16.318 46.682 1.00 3	8.95
	ATOM	92 CG PRO 212 36.409 17.156 47.604 1.00 4	12.00
	<b>ATOM</b>	93 C PRO 212 35.635 13.865 46.096 1.00 38	3.78
	ATOM	94 O PRO 212 36.546 13.714 45.280 1.00 34	4.64
	ATOM	95 N GLU 213 34.517 13.153 46.077 1.00 4	0.31
40	ATOM	96 CA GLU 213 34.256 12.160 45.049 1.00	
	ATOM	97 CB GLU 213 33.722 10.873 45.684 1.00 4	
	ATOM	98 CG GLU 213 34.616 10.344 46.800 1.00	
	ATOM	99 CD GLU 213 34.404 8.870 47.088 1.00 5	
	ATOM	100 OE1 GLU 213 33.240 8.416 47.072 1.00	
45	ATOM	101 OE2 GLU 213 35.402 8.167 47.353 1.00	
	ATOM	102 C GLU 213 33.234 12.796 44.083 1.00 4	
	ATOM	103 O GLU 213 32.703 13.876 44.368 1.00 4	
	ATOM	104 N PRO 214 32.953 12.154 42.933 1.00 4	
	ATOM	105 CD PRO 214 33.459 10.884 42.391 1.00	
50	ATOM	106 CA PRO 214 31.995 12.737 41.982 1.00	47.52

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            114 OG1 THR 215
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                               28.326 13.192 40.465 1.00 53.48
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            117 O THR 215
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             120 CB ASP 216
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                               24.150 12.941 41.270 1.00 70.57
             121 CG ASP 216
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                               23.902 12.257 42.595 1.00 78.07
             122 OD1 ASP 216
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     ATOM
             123 OD2 ASP 216
                                23.572 12.962 43.571 1.00 86.55
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                               25.706 13.277 39.356 1.00 58.42
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            125 O ASP 216
             126 N GLU 217
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            127 CA GLU 217
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     ATOM
            128 CB GLU 217
                                25.861 16.682 37.906 1.00 51.02
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            129 C GLU 217
                               27.211 14.692 37.195 1.00 53.55
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            130 O GLU 217
                               27.239 14.301 36.027 1.00 54.33
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            131 N GLU 218
                               28.290 14.726 37.975 1.00 49.20
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     ATOM
            132 CA GLU 218
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    ATOM
     ATOM
            133 CB GLU 218
                                30.674 14.601 38.530 1.00 43.43
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            134 CG GLU 218
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            135 CD GLU 218
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            137 OE2 GLU 218
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            139 O GLU 218
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                               28.892 10.572 37.539 1.00 46.97
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            142 CB TRP 219
                               28.183 9.762 38.630 1.00 48.42
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            144 CD2 TRP 219
                                30.167 8.572 39.879 1.00 55.24
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                                30.659 8.610 41.201 1.00 53.67
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                                30.795 7.745 38.938 1.00 54.55
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                                31.771 7.846 41.622 1.00 52.54
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             150 CZ3 TRP 219
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            151 CH2 TRP 219
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            153 O TRP 219
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            154 N GLU 220
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    ATOM
            156 CB GLU 220
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            158 CD GLU 220
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            159 OE1 GLU 220
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            160 OE2 GLU 220
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            161 C GLU 220
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            162 O GLU 220
                               27.399 10.899 32.443 1.00 49.94
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            163 N LEU 221
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                               29.868 14.317 33.155 1.00 39.21
            165 CB LEU 221
     ATOM
                                30.945 14.949 32.261 1.00 36.34
            166 CG LEU 221
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                                30.339 15.351 30.922 1.00 36.93
            167 CD1 LEU 221
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                                31.535 16.164 32.949 1.00 24.18
            168 CD2 LEU 221
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                               30.234 11.928 32.505 1.00 43.46
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                               30.618 11.621 31.375 1.00 45.25
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            170 O LEU 221
            171 N ILE 222
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            172 CA ILE 222
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            173 CB ILE 222
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            174 CG2 ILE 222
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            175 CG1 ILE 222
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                               33.111 10.646 37.199 1.00 34.85
            176 CD1 ILE 222
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            177 C ILE 222
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                              31.877 8.427 32.070 1.00 31.90
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            179 N LYS 223
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            180 CA LYS 223
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            181 CB LYS 223
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            182 CG LYS 223
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            183 CD LYS 223
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            190 CB THR 224
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            194 O THR 224
                               31.303 10.123 28.833 1.00 38.02
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            195 N VAL 225
            196 CA VAL 225
                                32.680 10.117 28.355 1.00 38.12
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            197 CB VAL 225
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                                34.960 11.162 28.632 1.00 36.77
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            198 CG1 VAL 225
            199 CG2 VAL 225
                                32.910 12.361 29.406 1.00 41.76
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            200 C VAL 225
                               33.291 8.724 28.302 1.00 37.52
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            201 O VAL 225
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            202 N THR 226
                               33.002 7.904 29.310 1.00 34.02
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     ATOM
            203 CA THR 226
                                33.542 6.552 29.350 1.00 34.67
                                33.237 5.857 30.707 1.00 30.56
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            205 OG1 THR 226
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            206 CG2 THR 226
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            208 O THR 226
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            210 CA GLU 227
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                               29.434 5.296 27.111 1.00 38.06
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                              31.466 5.409 25.651 1.00 37.69
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           213 O GLU 227
                              31.713 4.544 24.805 1.00 40.94
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           214 N ALA 228
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                               32.156 7.236 24.177 1.00 32.48
    ATOM 215 CA ALA 228
    ATOM 216 CB ALA 228
                               32.285 8.746 24.256 1.00 28.25
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           219 N HIS 229
                             34.404 6.611 24.843 1.00 33.58
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            220 CA HIS 229
                              35.724 6.029 24.669 1.00 32.97
    ATOM
            221 CB HIS 229
                              36.579 6.263 25.921 1.00 33.69
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            222 CG HIS 229
                              37.857 5.489 25.934 1.00 28.39
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            223 CD2 HIS 229
                               38.338 4.576 26.811 1.00 28.83
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            224 ND1 HIS 229
                               38.804 5.593 24.937 1.00 30.47
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           225 CE1 HIS 229
                              39.812 4.779 25.193 1.00 26.95
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                               39.556 4.147 26.332 1.00 31.27
           226 NE2 HIS 229
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    ATOM
           227 C HIS 229
                             35.653 4.536 24.371 1.00 38.40
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                             36.227 4.071 23.383 1.00 41.49
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    ATOM 231 CB VAL 230
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    ATOM 232 CG1 VAL 230
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           233 CG2 VAL 230
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                              34.219 1.934 23.700 1.00 44.28
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           234 C VAL 230
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           235 O VAL 230
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                               32.580 2.403 21.961 1.00 47.84
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            237 CA ALA 231
    ATOM
           238 CB ALA 231
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           240 O ALA 231
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    ATOM 243 CB THR 232
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    ATOM 244 OG1 THR 232
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    ATOM 245 CG2 THR 232
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            249 CA ASN 233
                               38.218 2.101 21.368 1.00 58.62
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            250 CB ASN 233
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            251 CG ASN 233
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                                40.843 2.401 22.724 1.00 65.50
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           252 OD1 ASN 233
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           253 ND2 ASN 233
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           254 C ASN 233
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             263 CB GLN 235
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            276 CB SER 237
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            278 O SER 237
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            279 N HIS 238
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             281 CB HIS 238
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            297 O TRP 239
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             310 CG ASN 241
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WO 99/26966

PCT/US98/25296

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     ATOM
                                40.448 11.777 18.076 1.00 28.64
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     ATOM
                                41.448 9.577 17.487 1.00 28.28
            586 CG2 VAL 283
    ATOM
            587 C VAL 283
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            588 O VAL 283
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                               37.403 10.945 16.028 1.00 30.96
            589 N VAL 284
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            590 CA VAL 284
                                36.293 11.838 15.694 1.00 29.14
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                                34.990 12.985 13.868 1.00 24.21
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            594 C VAL 284
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            606 CB PHE 286
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            607 CG PHE 286
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            609 CD2 PHE 286
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            631 CB LYS 289
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            635 NZ LYS 289
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            637 O LYS 289
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            643 CD2 LEU 290
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                               28.279 15.500 24.137 1.00 40.27
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     ATOM
            657 SD MET 292
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            659 C MET 292
660 O MET 292
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            669 CZ PHE 293
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            671 O PHE 293
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            675 SG CYS 294
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            676 C CYS 294
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     ATOM
            677 O CYS 294
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            678 N GLU 295
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            679 CA GLU 295
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            681 CG GLU 295
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	ATOM	684 OE2 GLU 295	24.017 22.029 22.199 1.00 85.30
	<b>ATOM</b>	685 C GLU 295	28.484 23.589 20.515 1.00 48.54
5	ATOM	686 O GLU 295	28.170 24.777 20.537 1.00 49.82
_	ATOM	687 N LEU 296	29.637 23.149 20.030 1.00 43.79
	ATOM	688 CA LEU 296	30,629 24.066 19.476 1.00 45.42
	ATOM	689 CB LEU 296	32.040 23.541 19.771 1.00 41.04
	ATOM	690 CG LEU 296	32.416 23.394 21.252 1.00 42.74
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- 0	ATOM	692 CD2 LEU 296	32.406 24.755 21.945 1.00 39.44
	ATOM	693 C LEU 296	30.448 24.239 17.968 1.00 45.56
	ATOM	694 O LEU 296	29.966 23.333 17.278 1.00 43.07
	ATOM	695 N PRO 297	30.823 25.414 17.436 1.00 46.99
15	ATOM	696 CD PRO 297	31.372 26.613 18.083 1.00 47.12
	ATOM	697 CA PRO 297	30.689 25.650 15.998 1.00 49.61
	ATOM	698 CB PRO 297	31.106 27.118 15.861 1.00 49.91
	ATOM	699 CG PRO 297	30,757 27.693 17.230 1.00 51.28
	ATOM	700 C PRO 297	31.600 24.717 15.202 1.00 49.59
20	ATOM	701 O PRO 297	32.727 24.446 15.615 1.00 51.66
	ATOM	702 N CYS 298	31.093 24.227 14.075 1.00 51.02
	<b>ATOM</b>	703 CA CYS 298	31.817 23.322 13.158 1.00 52.86
	ATOM	704 CB CYS 298	31.100 23.260 11.804 1.00 54.57
	ATOM	705 SG CYS 298	31.935 24.249 10.470 1.00 67.87
25	ATOM	706 C CYS 298	33.269 23.797 12.974 1.00 48.51
	<b>ATOM</b>	707 O CYS 298	34.197 22.991 12.819 1.00 49.58
	ATOM	708 N GLU 299	33.464 25.113 13.019 1.00 44.17
	ATOM	709 CA GLU 299	34.797 25.692 12.890 1.00 47.57
	ATOM	710 CB GLU 299	34.741 27.227 12.912 1.00 49.92
30	ATOM	711 CG GLU 299	34.001 27.871 11.747 1.00 59.30
	ATOM	712 CD GLU 299	32.489 27.763 11.848 1.00 63.80
	ATOM	713 OE1 GLU 299	31.805 28.162 10.882 1.00 69.03
	ATOM	714 OE2 GLU 299	31.979 27.297 12.889 1.00 67.10
	ATOM	715 C GLU 299	35.698 25.213 14.031 1.00 46.57
35	ATOM	716 O GLU 299	36.772 24.659 13.787 1.00 44.65
	ATOM	717 N ASP 300	35.263 25.432 15.274 1.00 45.17
	ATOM	718 CA ASP 300	36.046 25.008 16.433 1.00 43.32
	ATOM	719 CB ASP 300	35.442 25.517 17.747 1.00 37.38
	ATOM	720 CG ASP 300	35.567 27.016 17.910 1.00 36.23
40	ATOM	721 OD1 ASP 300	36.486 27.613 17.313 1.00 35.87
	ATOM	722 OD2 ASP 300	34.769 27.601 18.669 1.00 40.14
	ATOM	723 C ASP 300	36.174 23.495 16.513 1.00 42.81
	ATOM	724 O ASP 300	37.193 22.979 16.974 1.00 46.02
4.5	ATOM	725 N GLN 301	35.139 22.788 16.066 1.00 38.60
45	ATOM	726 CA GLN 301	35.151 21.334 16.086 1.00 40.00
	ATOM	727 CB GLN 301	33.815 20.783 15.576 1.00 38.59
	ATOM	728 CG GLN 301	32.608 21.334 16.317 1.00 40.26
	ATOM	729 CD GLN 301	31.311 20.696 15.869 1.00 44.15
60	ATOM	730 OE1 GLN 301	31.074 20.527 14.673 1.00 45.73
50	ATOM	731 NE2 GLN 301	30.450 20.363 16.824 1.00 46.13

	ATOM	732 C GLN 301	36.298 20.807 15.227 1.00 41.64
	ATOM	733 O GLN 301	36.975 19.850 15.601 1.00 45.02
	ATOM	734 N ILE 302	36.523 21.441 14.077 1.00 41.01
	ATOM	735 CA ILE 302	37.607 21.029 13.189 1.00 40.23
5	ATOM	736 CB ILE 302	37.580 21.798 11.825 1.00 39.52
	<b>ATOM</b>	737 CG2 ILE 302	38.724 21.308 10.931 1.00 31.98
	<b>ATOM</b>	738 CG1 ILE 302	36.230 21.607 11.119 1.00 40.77
	ATOM	739 CD1 ILE 302	35.895 20.166 10.733 1.00 45.43
	<b>ATOM</b>	740 C ILE 302	38.948 21.322 13.869 1.00 38.58
10	ATOM	741 O ILE 302	39.811 20.452 13.938 1.00 40.81
	ATOM	742 N ILE 303	39.110 22.547 14.364 1.00 37.50
	ATOM	743 CA ILE 303	40.343 22.958 15.030 1.00 39.33
	ATOM	744 CB ILE 303	40.263 24.442 15.501 1.00 39.06
	ATOM	745 CG2 ILE 303	41.525 24.822 16.279 1.00 36.19
15	ATOM	746 CG1 ILE 303	40.103 25.358 14.280 1.00 40.15
	ATOM	747 CD1 ILE 303	39.972 26.846 14.602 1.00 36.93
	ATOM	748 C ILE 303	40.676 22.061 16.222 1.00 36.49
	ATOM	749 O ILE 303	41.818 21.623 16.378 1.00 36.58
	ATOM	750 N LEU 304	39.674 21.788 17.057 1.00 32.91
20	ATOM	751 CA LEU 304	39.851 20.940 18.234 1.00 27.55
	<b>ATOM</b>	752 CB LEU 304	38.546 20.875 19.026 1.00 22.35
	ATOM	753 CG LEU 304	38.472 21.629 20.361 1.00 26.88
	ATOM	754 CD1 LEU 304	39.096 22.998 20.275 1.00 24.82
	ATOM	755 CD2 LEU 304	37.024 21.728 20.787 1.00 23.69
25	ATOM	756 C LEU 304	40.313 19.534 17.855 1.00 28.05
	ATOM	757 O LEU 304	41.277 19.013 18.429 1.00 24.68
	ATOM	758 N LEU 305	39.637 18.929 16.882 1.00 26.34
	ATOM	759 CA LEU 305	39.997 17.588 16.436 1.00 30.91
	ATOM	760 CB LEU 305	38.937 17.055 15.466 1.00 32.50
30	ATOM	761 CG LEU 305	37.585 16.757 16.132 1.00 33.36
	ATOM	762 CD1 LEU 305	36.557 16.439 15.079 1.00 33.87
	ATOM	763 CD2 LEU 305	37.733 15.581 17.101 1.00 31.72
	ATOM	764 C LEU 305	41.381 17.523 15.796 1.00 29.76
	ATOM	765 O LEU 305	42.109 16.553 15.990 1.00 29.33
35	ATOM	766 N LYS 306	41.754 18.554 15.048 1.00 29.72
	ATOM	767 CA LYS 306	43.065 18.569 14.409 1.00 34.28
	ATOM	768 CB LYS 306	43.122 19.673 13.345 1.00 35.98
	ATOM	769 CG LYS 306	42.140 19.465 12.206 1.00 43.35
	ATOM	770 CD LYS 306	42.195 20.583 11.170 1.00 51.50
40	ATOM	771 CE LYS 306	43.532 20.639 10.446 1.00 53.26
	ATOM	772 NZ LYS 306	43.522 21.702 9.409 1.00 59.61
	ATOM	773 C LYS 306	44.183 18.777 15.434 1.00 35.25
	ATOM	774 O LYS 306	45.312 18.332 15.231 1.00 33.95
	ATOM	775 N GLY 307	43.853 19.446 16.536 1.00 35.79
45	ATOM	776 CA GLY 307	44.836 19.700 17.576 1.00 34.59
	ATOM	777 C GLY 307	45.075 18.562 18.559 1.00 33.80
	ATOM	778 O GLY 307	46.200 18.360 19.008 1.00 31.59
	ATOM	779 N CYS 308	44.030 17.806 18.880 1.00 31.15
	ATOM	780 CA CYS 308	44.153 16.712 19.839 1.00 29.04
50	ATOM	781 CB CYS 308	42.929 16.667 20.750 1.00 27.59

WO 99/26966

PCT/US98/25296

	ATOM	782 SG CYS 308	41.452 15.974 19.941 1.00 30.50
	ATOM	783 C CYS 308	44,289 15.339 19.208 1.00 30.59
	ATOM	784 O CYS 308	44.609 14.374 19.899 1.00 33.77
	ATOM	785 N CYS 309	44.053 15.247 17.907 1.00 28.46
5	ATOM	786 CA CYS 309	44.099 13.961 17.219 1.00 30.10
J	ATOM	787 CB CYS 309	43.983 14.161 15.706 1.00 33.43
	ATOM	787 CB CTS 309	43.761 12.613 14.819 1.00 35.20
	ATOM	789 C CYS 309	45.301 13.071 17.524 1.00 27.72
	ATOM	790 O CYS 309	45.135 11.907 17.913 1.00 27.69
10	ATOM	791 N MET 310	46.508 13.594 17.339 1.00 26.15
10		791 N MET 310	47.700 12.798 17.605 1.00 26.06
	ATOM		48.928 13.439 16.951 1.00 25.32
	ATOM		50.207 12.648 17.132 1.00 23.32
	ATOM		
1.5	ATOM	795 SD MET 310	
15	ATOM	796 CE MET 310	51.674 10.307 16.934 1.00 28.50
	ATOM	797 C MET 310	47.941 12.612 19.113 1.00 25.94
	ATOM	798 O MET 310	48.592 11.653 19.526 1.00 28.09
	ATOM	799 N GLU 311	47.405 13.522 19.923 1.00 25.39
20	ATOM	800 CA GLU 311	47.560 13.445 21.370 1.00 27.03
20	ATOM	801 CB GLU 311	47.099 14.748 22.030 1.00 24.39
	ATOM	802 CG GLU 311	47.610 15.999 21.331 1.00 26.00
	ATOM	803 CD GLU 311	47.292 17.271 22.084 1.00 23.95
	ATOM	804 OE1 GLU 311	46.182 17.379 22.640 1.00 19.72
~-	ATOM	805 OE2 GLU 311	48.150 18.181 22.088 1.00 26.51
25	ATOM	806 C GLU 311	46.727 12.272 21.902 1.00 27.51
	ATOM	807 O GLU 311	47.152 11.552 22.807 1.00 29.67
	ATOM	808 N ILE 312	45.547 12.086 21.326 1.00 26.82
	ATOM	809 CA ILE 312	44.661 11.001 21.724 1.00 25.71
• •	ATOM	810 CB ILE 312	43.194 11.296 21.304 1.00 23.35
30	ATOM	811 CG2 ILE 312	42.301 10.068 21.583 1.00 20.27
	ATOM	812 CG1 ILE 312	42.690 12.534 22.062 1.00 20.88
	ATOM	813 CD1 ILE 312	41.244 12.961 21.755 1.00 18.15
	ATOM	814 C ILE 312	45.116 9.665 21.132 1.00 27.91
	ATOM	815 O ILE 312	45.064 8.628 21.804 1.00 28.96
35	ATOM	816 N MET 313	45.582 9.683 19.886 1.00 27.66
	ATOM	817 CA MET 313	46.045 8.447 19.257 1.00 30.18
	ATOM	818 CB MET 313	46.386 8.662 17.771 1.00 36.89
	ATOM	819 CG MET 313	45.186 8.938 16.861 1.00 37.95
	ATOM	820 SD MET 313	45.624 8.943 15.096 1.00 42.38
40	ATOM	821 CE MET 313	46.724 10.319 14.999 1.00 40.68
	ATOM	822 C MET 313	47.264 7.897 19.975 1.00 27.43
	ATOM	823 O MET 313	47.351 6.690 20.219 1.00 28.61
	ATOM	824 N SER 314	48.202 8.776 20.318 1.00 24.88
	ATOM	825 CA SER 314	49.416 8.352 21.011 1.00 27.98
45	ATOM	826 CB SER 314	50.420 9.511 21.118 1.00 29.64
	ATOM	827 OG SER 314	49.912 10.560 21.911 1.00 43.44
	ATOM	828 C SER 314	49.082 7.818 22.402 1.00 22.30
	<b>ATOM</b>	829 O SER 314	49.737 6.895 22.892 1.00 24.18
	<b>ATOM</b>	830 N LEU 315	48.070 8.395 23.039 1.00 23.99
50	ATOM	831 CA LEU 315	47.646 7.918 24.365 1.00 25.07

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46.580 8.842 24.965 1.00 19.11
            832 CB LEU 315
    ATOM
                               45.863 8.355 26.228 1.00 20.39
            833 CG LEU 315
    ATOM
                                46.872 8.076 27.362 1.00 18.92
            834 CD1 LEU 315
    ATOM
                                44.848 9.401 26.655 1.00 12.93
            835 CD2 LEU 315
    ATOM
                              47.070 6.518 24.222 1.00 24.53
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    ATOM
            836 C LEU 315
                              47.394 5.615 24.992 1.00 26.32
    ATOM
            837 O LEU 315
            838 N ARG 316
                              46.212 6.338 23.220 1.00 28.18
    ATOM
                               45.595 5.041 22.978 1.00 27.54
            839 CA ARG 316
    ATOM
                               44.575 5.155 21.848 1.00 27.39
    ATOM
            840 CB ARG 316
                               43.340 5.929 22.253 1.00 22.00
    ATOM
            841 CG ARG 316
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    ATOM
            842 CD ARG 316
                               42.291 5.902 21.172 1.00 18.78
    ATOM
            843 NE ARG 316
                               40.975 6.205 21.719 1.00 26.57
            844 CZ ARG 316
                               39.852 6.224 21.014 1.00 30.81
    ATOM
            845 NH1 ARG 316
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    ATOM
            846 NH2 ARG 316
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15
            847 C ARG 316
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    ATOM
            849 N ALA 317
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    ATOM 850 CA ALA 317
    ATOM 851 CB ALA 317
                               49.674 3.904 20.643 1.00 22.93
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            852 C ALA 317
                              49.591 3.115 23.002 1.00 28.35
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            853 O ALA 317
                              49.968 1.979 23.312 1.00 32.10
    ATOM
            854 N ALA 318
                              49.863 4.197 23.727 1.00 29.12
     ATOM
                               50.655 4.123 24.953 1.00 27.50
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            855 CA ALA 318
                               50.854 5.518 25.522 1.00 28.39
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            856 CB ALA 318
                               50.053 3.215 26.013 1.00 28.10
    ATOM 857 C ALA 318
            858 O ALA 318
859 N VAL 319
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                               48.730 3.245 26.165 1.00 29.16
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     ATOM 860 CA VAL 319
                               48.082 2.414 27.176 1.00 35.24
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                               46.759 4.324 28.136 1.00 29.96
            862 CG1 VAL 319
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                               45.773 2.936 26.322 1.00 31.70
    ATOM 863 CG2 VAL 319
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    ATOM 864 C VAL 319
    ATOM 865 O VAL 319
ATOM 866 N ARG 320
                               47.448 0.129 27.515 1.00 42.70
                               48.460 0.644 25.565 1.00 38.64
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                               47.764 -0.751 23.674 1.00 37.26
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    ATOM
                               46.258 -0.655 23.720 1.00 43.12
            869 CG ARG 320
    ATOM
            870 CD ARG 320
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                               44.260 -0.446 22.286 1.00 54.71
            871 NE ARG 320
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            872 CZ ARG 320
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     ATOM
                                44.119 0.713 20.300 1.00 49.08
            873 NH1 ARG 320
     ATOM
            874 NH2 ARG 320
                                42.206 -0.058 21.326 1.00 59.59
     ATOM
                               49.852 -1.247 24.930 1.00 42.14
     ATOM
            875 C ARG 320
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     ATOM 876 O ARG 320
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                               50.712 -0.772 25.822 1.00 42.04
     ATOM 877 N TYR 321
                               52.098 -1.202 25.852 1.00 42.70
     ATOM 878 CA TYR 321
           879 CB TYR 321
                               52.971 -0.133 26.529 1.00 38.01
     ATOM
    ATOM 880 CG TYR 321
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    ATOM 881 CD1 TYR 321
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             883 CD2 TYR 321
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     ATOM
             884 CE2 TYR 321
                                 56.194 -1.411 28.207 1.00 32.69
     ATOM
             885 CZ TYR 321
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     ATOM
             886 OH TYR 321
                                 58.289 -2.158 27.288 1.00 39.48
     ATOM
                               52.189 -2.515 26.629 1.00 45.51
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             887 C TYR 321
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             888 O TYR 321
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     ATOM
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                                53.129 -4.764 26.753 1.00 45.86
     ATOM
            891 CB ASP 322
                                52.697 -5.899 25.816 1.00 46.64
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     ATOM
            892 C ASP 322
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     ATOM
            893 O ASP 322
                               55.434 -5.109 26.214 1.00 45.38
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             894 N PRO 323
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     ATOM
             895 CD PRO 323
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             896 CA PRO 323
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             897 CB PRO 323
                                56.293 -4.667 30.308 1.00 43.95
     ATOM
     ATOM
             898 CG PRO 323
                                54.926 -5.223 30.655 1.00 43.93
             899 C PRO 323
     ATOM
                               56.993 -6.285 28.478 1.00 48.34
             900 O PRO 323
                               58.217 -6.407 28.379 1.00 50.84
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             901 N GLU 324
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                                56.149 -7.301 28.315 1.00 52.39
            902 CA GLU 324
                                56.621 -8.646 28.005 1.00 55.85
    ATOM
            903 CB GLU 324
                                55.453 -9.633 28.048 1.00 55.54
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            904 C GLU 324
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                                57.283 -8.670 26.632 1.00 54.94
            905 O GLU 324
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                                58.460 -9.013 26.502 1.00 59.81
            906 N SER 325
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     ATOM
            907 CA SER 325
     ATOM
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             908 CB SER 325
                                55.889 -8.613 23.279 1.00 48.23
     ATOM
             909 OG SER 325
                                54.788 -7.749 23.471 1.00 48.71
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                               58.194 -6.743 22.808 1.00 52.19
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             911 O SER 325
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             912 N GLU 326
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            913 CA GLU 326
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    ATOM
            914 CB GLU 326
                                59.469 -4.562 24.587 1.00 42.74
     ATOM
            915 CG GLU 326
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     ATOM
            916 CD GLU 326
                                61.565 -5.067 25.907 1.00 56.34
     ATOM
            917 OE1 GLU 326
                                 62.139 -5.407 26.966 1.00 59.31
            918 OE2 GLU 326
                                 62.178 -4.774 24.856 1.00 55.74
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     ATOM
            919 C GLU 326
                                57.397 -3.993 23.291 1.00 40.23
            920 O GLU 326
                                58.145 -3.474 22.465 1.00 40.44
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            921 N THR 327
                                56.080 -4.079 23.127 1.00 35.90
     ATOM
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             922 CA THR 327
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     ATOM
            923 CB THR 327
                                54.983 -4.717 21.008 1.00 37.63
             924 OG1 THR 327
    ATOM
                                 53.994 -5.503 21.674 1.00 38.12
                                 56.165 -5.609 20.635 1.00 39.90
     ATOM
             925 CG2 THR 327
                               54.170 -2.780 22.282 1.00 39.49
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            926 C THR 327
     ATOM
            927 O THR 327
     ATOM
                                53.603 -2.930 23.364 1.00 40.50
     ATOM
            928 N LEU 328
                               53.758 -1.933 21.347 1.00 36.64
     ATOM
            929 CA LEU 328
                                52.544 -1.136 21.480 1.00 37.73
            930 CB LEU 328
                                52.791 0.340 21.127 1.00 37.78
     ATOM
                                53.667 1.257 21.982 1.00 36.26
            931 CG LEU 328
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     ATOM
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     ATOM
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             933 CD2 LEU 328
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            934 C LEU 328
     ATOM
                               52.083 -2.233 19.410 1.00 34.96
     ATOM
            935 O LEU 328
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            936 N THR 329
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     ATOM
            937 CA THR 329
            938 CB THR 329
                                48.401 -3.176 20.349 1.00 42.67
     ATOM
            939 OG1 THR 329
                                 49.156 -4.271 20.896 1.00 42.52
     ATOM
            940 CG2 THR 329
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                                48,591 -1.034 19.058 1.00 44.31
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     ATOM
            941 C THR 329
     ATOM
            942 O THR 329
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     ATOM
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             944 CA LEU 330
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                                49.056 0.545 15.766 1.00 44.66
             945 CB LEU 330
     ATOM
     ATOM
             946 CG LEU 330
                                50.329 1.393 15.951 1.00 51.06
15
                                 51.195 0.890 17.095 1.00 48.58
     ATOM
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            948 CD2 LEU 330
                                 51.107 1.387 14.638 1.00 45.18
     ATOM
            949 C LEU 330
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     ATOM
            950 O LEU 330
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            951 N ASN 331
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     ATOM
            952 CA ASN 331
                                44.450 0.363 16.257 1.00 54.41
     ATOM
            953 CB ASN 331
                                44.370 0.353 14.731 1.00 54.94
     ATOM
            954 CG ASN 331
                                42.970 0.603 14.219 1.00 60.35
     ATOM
                                 42.375 1.642 14.501 1.00 61.84
     ATOM
            955 OD1 ASN 331
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            956 ND2 ASN 331
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     ATOM
     ATOM
            957 C ASN 331
                               43.940 -0.963 16.836 1.00 58.00
                                42.985 -1.557 16.328 1.00 60.17
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            958 O ASN 331
            959 N GLY 332
960 CA GLY 332
                                44.590 -1.414 17.908 1.00 58.45
     ATOM
                                44.215 -2.658 18.556 1.00 58.55
     ATOM
            961 C GLY 332
                                44.408 -3.880 17.680 1.00 59.79
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     ATOM
            962 O GLY 332
                                43.892 -4.953 17.993 1.00 61.32
     ATOM
                                45,165 -3.725 16.597 1.00 60.28
            963 N GLU 333
     ATOM
     ATOM
            964 CA GLU 333
                                45.408 -4.821 15.659 1.00 59.13
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     ATOM
            966 CG GLU 333
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     ATOM
            967 CD GLU 333
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     ATOM
                                 43.374 -2.740 12.359 1.00 79.98
            968 OE1 GLU 333
     ATOM
            969 OE2 GLU 333
                                 41.942 -4.412 12.392 1.00 83.81
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            970 C GLU 333
                                46,881 -5.146 15.452 1.00 57.18
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     ATOM
            971 O GLU 333
                                47.291 -6.301 15.545 1.00 57.50
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            972 N MET 334
                                47.663 -4.112 15.166 1.00 55.20
     ATOM
            973 CA MET 334
974 CB MET 334
                                 49.085 -4.245 14.873 1.00 50.85
     ATOM
                                 49.416 -3.334 13.687 1.00 48.70
     ATOM
                                 50.844 -3.412 13.181 1.00 45.39
     ATOM
            975 CG MET 334
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            976 SD MET 334
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     ATOM
                                49.908 -2.477 10.749 1.00 45.25
     ATOM
            977 CE MET 334
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            978 C MET 334
     ATOM
            979 O MET 334
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     ATOM
            980 N ALA 335
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     ATOM
                                51.769 -4.787 17.527 1.00 48.98
            981 CA ALA 335
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     ATOM
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     ATOM
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            984 O ALA 335
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                               53.718 -3.336 17.307 1.00 46.62
            985 N VAL 336
     ATOM
    ATOM
                               55.016 -2.926 16.783 1.00 42.35
            986 CA VAL 336
                               54.876 -1.687 15.877 1.00 42.41
     ATOM
           987 CB VAL 336
     ATOM 988 CG1 VAL 336
                                53.963 -2.004 14.707 1.00 42.00
     ATOM 989 CG2 VAL 336
                                54.313 -0.506 16.676 1.00 40.32
                               56.023 -2.608 17.883 1.00 45.33
     ATOM 990 C VAL 336
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     ATOM 991 O VAL 336
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            992 N THR 337
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    ATOM
            993 CA THR 337
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            994 CB THR 337
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     ATOM
            995 OG1 THR 337
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     ATOM
    ATOM 996 CG2 THR 337
                                59.253 -4.734 18.392 1.00 40.38
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    ATOM 997 C THR 337
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     ATOM 998 O THR 337
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     ATOM 999 N ARG 338
                               59.655 -0.489 19.268 1.00 37.61
     ATOM 1000 CA ARG 338
                                60.171 0.876 19.268 1.00 38.68
                                61.177 1.041 20.424 1.00 35.95
     ATOM 1001 CB ARG 338
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     ATOM 1002 CG ARG 338
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    ATOM 1003 CD ARG 338
                                62.114 2.277 23.035 1.00 37.42
     ATOM 1004 NE ARG 338
     ATOM 1005 CZ ARG 338
                                61.858 3.256 23.902 1.00 30.20
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    ATOM 1006 NH1 ARG 338
                                62.224 4.501 23.636 1.00 27.98
    ATOM 1007 NH2 ARG 338
                                61.213 2.992 25.025 1.00 27.40
    ATOM 1008 C ARG 338
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     ATOM 1009 O ARG 338
                               60.529 2.142 17.251 1.00 34.12
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    ATOM 1011 CA GLY 339
                                62.475 0.416 16.282 1.00 41.35
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    ATOM 1012 C GLY 339
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    ATOM 1013 O GLY 339
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    ATOM 1014 N GLN 340
                               60.594 -0.414 14.982 1.00 38.58
     ATOM 1015 CA GLN 340
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    ATOM 1016 CB GLN 340
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                                59.450 -2.995 13.944 1.00 41.10
    ATOM 1017 CG GLN 340
    ATOM 1018 CD GLN 340
                                58.468 -4.144 13.890 1.00 48.84
    ATOM 1019 OE1 GLN 340
                                57.529 -4.208 14.679 1.00 50.53
    ATOM 1020 NE2 GLN 340
                                58.685 -5.068 12.959 1.00 54.25
    ATOM 1021 C GLN 340
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                               58.884 0.822 13.679 1.00 41.50
    ATOM 1022 O GLN 340
                               58.725 1.342 12.576 1.00 42.72
    ATOM 1023 N LEU 341
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                                57.546 2.532 14.775 1.00 38.10
    ATOM 1024 CA LEU 341
    ATOM 1025 CB LEU 341
                                56.868 2.740 16.133 1.00 36.66
45
    ATOM 1026 CG LEU 341
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    ATOM 1027 CD1 LEU 341
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                                55.389 3.989 17.700 1.00 40.95
    ATOM 1028 CD2 LEU 341
                               58.404 3.743 14.423 1.00 36.37
    ATOM 1029 C LEU 341
    ATOM 1030 O LEU 341
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    ATOM 1031 N LYS 342
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                                61.801 4.687 15.582 1.00 34.97
     ATOM 1033 CB LYS 342
                                62.764 5.863 15.519 1.00 40.00
     ATOM 1034 CG LYS 342
     ATOM 1035 CD LYS 342
                                63.868 5.739 16.555 1.00 34.48
     ATOM 1036 CE LYS 342
                                64.709 7.001 16.596 1.00 37.54
     ATOM 1037 NZ LYS 342
                                65.716 6.972 17.689 1.00 42.32
     ATOM 1038 C LYS 342
                               60.928 4.970 13.235 1.00 38.29
     ATOM 1039 O LYS 342
                               60.621 5.963 12.569 1.00 36.23
     ATOM 1040 N ASN 343
                               61.585 3.932 12.721 1.00 39.25
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                                62.014 3.903 11.328 1.00 40.19
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     ATOM 1042 CB ASN 343
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     ATOM 1043 CG ASN 343
                                63.937 2.429 12.027 1.00 39.22
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     ATOM 1044 OD1 ASN 343
     ATOM 1045 ND2 ASN 343
                                 64.125 1.197 12.471 1.00 42.19
     ATOM 1046 C ASN 343
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     ATOM 1047 O ASN 343
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     ATOM 1049 CA GLY 344
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     ATOM 1050 C GLY 344
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     ATOM 1051 O GLY 344
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     ATOM 1052 N GLY 345
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     ATOM 1053 CA GLY 345
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     ATOM 1054 C GLY 345
                               58.050 8.444 11.300 1.00 38.64
     ATOM 1055 O GLY 345
                               57.902 9.652 11.116 1.00 38.14
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     ATOM 1057 CA LEU 346
                                57.904 8.761 13.692 1.00 36.05
     ATOM 1058 CB LEU 346
                                57.039 8.048 14.738 1.00 35.72
     ATOM 1059 CG LEU 346
                                55.561 7.864 14.371 1.00 34.89
     ATOM 1060 CD1 LEU 346
                                54.850 7.132 15.494 1.00 44.09
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     ATOM 1061 CD2 LEU 346
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     ATOM 1062 C LEU 346
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     ATOM 1063 O LEU 346
                               59.171 10.257 15.066 1.00 35.58
     ATOM 1064 N GLY 347
                               60.299 8.595 14.067 1.00 30.47
     ATOM 1065 CA GLY 347
                                61.559 9.017 14.661 1.00 33.01
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    ATOM 1066 C GLY 347
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    ATOM 1067 O GLY 347
ATOM 1068 N VAL 348
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    ATOM 1069 CA VAL 348
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    ATOM 1070 CB VAL 348
                                62.843 11.612 18.620 1.00 31.66
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    ATOM 1071 CG1 VAL 348
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    ATOM 1072 CG2 VAL 348
                                63.080 11.651 20.118 1.00 24.77
    ATOM 1073 C VAL 348
                               60.683 10.273 18.855 1.00 33.84
    ATOM 1074 O VAL 348
                               60.546 10.034 20.050 1.00 29.99
    ATOM 1075 N VAL 349
                               59.649 10.518 18.049 1.00 33.31
    ATOM 1076 CA VAL 349
45
                                58.270 10.495 18.538 1.00 32.23
    ATOM 1077 CB VAL 349
                                57.279 10.911 17.415 1.00 32.59
    ATOM 1078 CG1 VAL 349
                                55.837 10.678 17.838 1.00 33.68
    ATOM 1079 CG2 VAL 349
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    ATOM 1080 C VAL 349
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    ATOM 1081 O VAL 349
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58.551 8.081 18.444 1.00 32.81
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    ATOM 1083 CA SER 350
                               59.041 5.746 17.904 1.00 24.95
    ATOM 1084 CB SER 350
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    ATOM 1085 OG SER 350
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    ATOM 1086 C SER 350
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    ATOM 1087 O SER 350
                               60.055 7.007 20.546 1.00 28.60
    ATOM 1088 N ASP 351
    ATOM 1089 CA ASP 351
                               60.652 6.863 21.867 1.00 29.82
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    ATOM 1090 CB ASP 351
    ATOM 1091 CG ASP 351
                               63.030 6.806 21.000 1.00 30.22
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    ATOM 1092 OD1 ASP 351
                                63.411 7.412 19.974 1.00 32.61
    ATOM 1093 OD2 ASP 351
                                63.422 5.661 21.301 1.00 30.02
    ATOM 1094 C ASP 351
                               59.785 7.548 22.913 1.00 30.63
                               59.632 7.055 24.027 1.00 29.54
    ATOM 1095 O ASP 351
                               59.222 8.692 22.537 1.00 25.33
    ATOM 1096 N ALA 352
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    ATOM 1097 CA ALA 352
                                58.390 9.464 23.432 1.00 28.59
                                58.011 10.798 22.788 1.00 20.95
    ATOM 1098 CB ALA 352
                               57.136 8.695 23.831 1.00 29.69
    ATOM 1099 C ALA 352
                               56,711 8,753 24,982 1,00 30,36
    ATOM 1100 O ALA 352
    ATOM 1101 N ILE 353
                              56.557 7.979 22.876 1.00 27.63
20
    ATOM 1102 CA ILE 353
                               55.345 7.227 23.129 1.00 27.55
    ATOM 1103 CB ILE 353
                               54.611 6.925 21.805 1.00 28.04
    ATOM 1104 CG2 ILE 353
                               53.329 6.111 22.065 1.00 23.68
                               54.269 8.251 21.119 1.00 27.33
    ATOM 1105 CG1 ILE 353
                               53.637 8.105 19.734 1.00 26.23
25
    ATOM 1106 CD1 ILE 353
                              55.631 5.943 23.901 1.00 30.88
    ATOM 1107 C ILE 353
    ATOM 1108 O ILE 353
                              54.880 5.597 24.814 1.00 31.22
    ATOM 1109 N PHE 354
                               56.710 5.240 23.549 1.00 29.86
    ATOM 1110 CA PHE 354
                               57.056 4.022 24.275 1.00 31.08
                               58.227 3.274 23.619 1.00 28.80
    ATOM 1111 CB PHE 354
30
                                57.799 2.322 22.523 1.00 28.80
    ATOM 1112 CG PHE 354
                                57.330 2.804 21.292 1.00 30.96
    ATOM 1113 CD1 PHE 354
    ATOM 1114 CD2 PHE 354
                                57.811 0.939 22.749 1.00 29.45
    ATOM 1115 CE1 PHE 354
                                56.864 1.909 20.281 1.00 27.12
    ATOM 1116 CE2 PHE 354
                                57.354 0.026 21.761 1.00 25.19
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    ATOM 1117 CZ PHE 354
                               56.879 0.518 20.521 1.00 28.09
    ATOM 1118 C PHE 354
                               57.398 4.349 25.721 1.00 29.17
                               57.001 3.625 26.631 1.00 32.62
    ATOM 1119 O PHE 354
     ATOM 1120 N ASP 355
                               58.133 5.438 25.925 1.00 23.86
    ATOM 1121 CA ASP 355
                               58.508 5.873 27.262 1.00 25.34
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     ATOM 1122 CB ASP 355
                               59.434 7.083 27.180 1.00 21.41
                                60.846 6.708 26.769 1.00 32.08
     ATOM 1123 CG ASP 355
     ATOM 1124 OD1 ASP 355
                                61.051 5.595 26.226 1.00 33.58
     ATOM 1125 OD2 ASP 355
                                61.756 7.534 26.970 1.00 33.20
                               57.254 6.211 28.062 1.00 27.86
45
     ATOM 1126 C ASP 355
                               57.167 5.916 29.252 1.00 32.42
     ATOM 1127 O ASP 355
     ATOM 1128 N LEU 356
                               56,276 6.821 27.401 1.00 26.84
     ATOM 1129 CA LEU 356
                                55.031 7.164 28.066 1.00 28.66
                                54.112 7.953 27.131 1.00 25.37
     ATOM 1130 CB LEU 356
                                52.787 8.427 27.742 1.00 27.61
    ATOM 1131 CG LEU 356
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53.056 9.452 28.842 1.00 25.43
     ATOM 1132 CD1 LEU 356
                                51.924 9.057 26.667 1.00 27.49
     ATOM 1133 CD2 LEU 356
                               54,334 5.875 28.473 1.00 30.44
     ATOM 1134 C LEU 356
                               53.873 5.743 29.601 1.00 31.55
     ATOM 1135 O LEU 356
                               54.266 4.928 27.536 1.00 32.69
    ATOM 1136 N GLY 357
                                53.621 3.652 27.787 1.00 29.87
     ATOM 1137 CA GLY 357
     ATOM 1138 C GLY 357
                               54.239 2.884 28.939 1.00 33.12
     ATOM 1139 O GLY 357
                               53.524 2.268 29.732 1.00 29.41
    ATOM 1140 N MET 358
                               55.570 2.911 29.026 1.00 33.31
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    ATOM 1141 CA MET 358
                                56.277 2.217 30.100 1.00 35.87
     ATOM 1142 CB MET 358
                                57.794 2.265 29.871 1.00 34.56
    ATOM 1143 CG MET 358
                                58.265 1.608 28.576 1.00 46.43
     ATOM 1144 SD MET 358
                                60.073 1.600 28.351 1.00 42.13
    ATOM 1145 CE MET 358
                                60.429 3.306 28.411 1.00 44.29
    ATOM 1146 C MET 358
                               55.948 2.884 31.434 1.00 33.26
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     ATOM 1147 O MET 358
                               55.802 2.222 32.453 1.00 36.39
                               55.825 4.202 31.398 1.00 33.31
     ATOM 1148 N SER 359
     ATOM 1149 CA SER 359
                               55.533 4.998 32.580 1.00 34.39
                               55.859 6.463 32.303 1.00 30.84
    ATOM 1150 CB SER 359
    ATOM 1151 OG SER 359
                              55.487 7.265 33.404 1.00 47.14
20
                              54.094 4.897 33.072 1.00 36.43
     ATOM 1152 C SER 359
                               53.833 5.073 34.260 1.00 35.46
    ATOM 1153 O SER 359
     ATOM 1154 N LEU 360
                               53.165 4.617 32.156 1.00 36.74
    ATOM 1155 CA LEU 360
                               51.750 4.519 32.493 1.00 35.44
25
    ATOM 1156 CB LEU 360
                               50.889 4.817 31.263 1.00 34.16
    ATOM 1157 CG LEU 360
                                50.896 6.263 30.751 1.00 34.59
     ATOM 1158 CD1 LEU 360
                                50.031 6.353 29.513 1.00 33.53
                                50.376 7.211 31.836 1.00 31.69
     ATOM 1159 CD2 LEU 360
     ATOM 1160 C LEU 360
                               51.324 3.192 33.088 1.00 38.72
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    ATOM 1161 O LEU 360
                               50.185 3.058 33.546 1.00 38.29
    ATOM 1162 N SER 361
                               52.227 2.214 33.080 1.00 40.96
     ATOM 1163 CA SER 361
                               51.938 0.897 33.636 1.00 45.67
                               53.131 -0.044 33.436 1.00 46.45
    ATOM 1164 CB SER 361
                               53.362 -0.296 32.061 1.00 51.81
     ATOM 1165 OG SER 361
                              51.628 1.004 35.124 1.00 44.49
    ATOM 1166 C SER 361
35
    ATOM 1167 O SER 361
                               50.724 0.337 35.630 1.00 46.67
    ATOM 1168 N SER 362
                               52.385 1.858 35.809 1.00 41.44
                               52.231 2.081 37.245 1.00 42.13
    ATOM 1169 CA SER 362
    ATOM 1170 CB SER 362
                               53.431 2.876 37.779 1.00 42.61
                               54.647 2.215 37.492 1.00 51.87
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    ATOM 1171 OG SER 362
                              50.951 2.832 37.610 1.00 38.41
    ATOM 1172 C SER 362
                               50.444 2.700 38.722 1.00 38.01
     ATOM 1173 O SER 362
                               50.443 3.631 36.672 1.00 34.55
    ATOM 1174 N PHE 363
    ATOM 1175 CA PHE 363
                               49.232 4.404 36.906 1.00 32.96
                               49.109 5.518 35.859 1.00 31.99
45
    ATOM 1176 CB PHE 363
                               50.093 6.659 36.058 1.00 29.97
    ATOM 1177 CG PHE 363
    ATOM 1178 CD1 PHE 363
                                49.667 7.872 36.594 1.00 30.61
                                51.445 6.501 35.731 1.00 32.02
     ATOM 1179 CD2 PHE 363
    ATOM 1180 CE1 PHE 363
                                50.579 8.940 36.803 1.00 33.67
                                52.376 7.552 35.934 1.00 30.91
50
    ATOM 1181 CE2 PHE 363
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	ATOM	1182 CZ PHE 363	51.938 8.777 36.473 1.00 29.33
	ATOM	1183 C PHE 363	47.973 3.554 36.916 1.00 30.52
	<b>ATOM</b>	1184 O PHE 363	46.971 3.947 37.491 1.00 32.19
	<b>ATOM</b>	1185 N ASN 364	48.036 2.384 36.283 1.00 33.51
5	ATOM	1186 CA ASN 364	46.894 1.471 36.216 1.00 38.03
	ATOM	1187 CB ASN 364	46.754 0.711 37.539 1.00 42.32
	ATOM	1188 CG ASN 364	47.824 -0.361 37.713 1.00 53.11
	ATOM	1189 OD1 ASN 364	47.815 -1.370 37.012 1.00 59.51
	ATOM	1190 ND2 ASN 364	48.751 -0.138 38.639 1.00 55.95
10	ATOM	1191 C ASN 364	45.574 2.161 35.871 1.00 31.89
	<b>ATOM</b>	1192 O ASN 364	44.587 2.027 36.588 1.00 30.28
	ATOM	1193 N LEU 365	45.561 2.883 34.751 1.00 27.62
	ATOM	1194 CA LEU 365	44.365 3.606 34.317 1.00 29.36
	ATOM	1195 CB LEU 365	44.738 4.627 33.240 1.00 27.54
15	ATOM	1196 CG LEU 365	45.826 5.659 33.576 1.00 38.91
	ATOM	1197 CD1 LEU 365	46.115 6.499 32.338 1.00 34.47
	<b>ATOM</b>	1198 CD2 LEU 365	45.394 6.546 34.743 1.00 34.24
	ATOM	1199 C LEU 365	43.264 2.691 33.774 1.00 26.23
	<b>ATOM</b>	1200 O LEU 365	43.546 1.648 33.197 1.00 27.06
20	ATOM	1201 N ASP 366	42.011 3.074 33.991 1.00 25.23
	ATOM	1202 CA ASP 366	40.892 2.307 33.462 1.00 26.07
	ATOM	1203 CB ASP 366	39.832 2.008 34.538 1.00 29.68
	ATOM	1204 CG ASP 366	39.337 3.253 35.261 1.00 35.74
	ATOM	1205 OD1 ASP 366	39.438 4.371 34.717 1.00 36.78
25	ATOM	1206 OD2 ASP 366	38.803 3.100 36.378 1.00 41.23
	ATOM	1207 C ASP 366	40.274 3.100 32.305 1.00 27.70
	ATOM	1208 O ASP 366	40.748 4.191 31.975 1.00 31.94
	ATOM	1209 N ASP 367	39.223 2.564 31.693 1.00 29.18
	ATOM	1210 CA ASP 367	38.594 3.233 30.560 1.00 32.72
30	ATOM	1211 CB ASP 367	37.428 2.395 30.018 1.00 38.04
	ATOM	1212 CG ASP 367	37.855 0.995 29.606 1.00 42.43
	ATOM	1213 OD1 ASP 367	38.913 0.852 28.956 1.00 35.95
	ATOM	1214 OD2 ASP 367	37.115 0.034 29.917 1.00 51.42
	ATOM	1215 C ASP 367	38.093 4.631 30.881 1.00 33.71
35	ATOM	1216 O ASP 367	38.059 5.506 30.013 1.00 38.30
	ATOM	1217 N THR 368	37.705 4.852 32.132 1.00 31.06
	ATOM	1218 CA THR 368	37.199 6.155 32.543 1.00 26.28
	ATOM	1219 CB THR 368	36.537 6.066 33.922 1.00 27.30
	ATOM	1220 OG1 THR 368	35.461 5.127 33.861 1.00 33.42
40	ATOM	1221 CG2 THR 368	36.003 7.423 34.355 1.00 25.16
	ATOM	1222 C THR 368	38.303 7.194 32.593 1.00 21.13
	ATOM	1223 O THR 368	38.133 8.314 32.104 1.00 23.17
	ATOM	1224 N GLU 369	39.431 6.816 33.179 1.00 21.32
	ATOM	1225 CA GLU 369	40.565 7.720 33.317 1.00 28.00
45	ATOM	1226 CB GLU 369	41.582 7.107 34.277 1.00 32.79
	ATOM	1227 CG GLU 369	40.944 6.804 35.619 1.00 36.29
	ATOM	1228 CD GLU 369	41.834 6.026 36.546 1.00 41.03
	ATOM	1229 OE1 GLU 369	42.361 4.967 36.123 1.00 42.05
	ATOM	1230 OE2 GLU 369	41.986 6.458 37.705 1.00 42.03
50	ATOM	1231 C GLU 369	41.201 8.047 31.970 1.00 25.57

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41.626 9.175 31.741 1.00 20.56
    ATOM 1232 O GLU 369
                               41.249 7.055 31.080 1.00 25.39
    ATOM 1233 N VAL 370
                                41.794 7.278 29.745 1.00 25.99
    ATOM 1234 CA VAL 370
                                42.005 5.936 28.977 1.00 26.15
    ATOM 1235 CB VAL 370
    ATOM 1236 CG1 VAL 370
                                42.450 6.216 27.539 1.00 27.65
    ATOM 1237 CG2 VAL 370
                                43.056 5.086 29.685 1.00 17.70
                               40.814 8.164 28.966 1.00 26.49
    ATOM 1238 C VAL 370
    ATOM 1239 O VAL 370
                               41.226 9.038 28.202 1.00 28.16
                               39.514 7.950 29.184 1.00 21.01
    ATOM 1240 N ALA 371
                                38.486 8.730 28.510 1.00 19.57
    ATOM 1241 CA ALA 371
10
                                37.116 8.136 28.783 1.00 18.62
    ATOM 1242 CB ALA 371
                               38,512 10,191 28,947 1,00 23,48
    ATOM 1243 C ALA 371
                               38.500 11.103 28.111 1.00 32.67
    ATOM 1244 O ALA 371
                               38,540 10,414 30.256 1.00 22.89
    ATOM 1245 N LEU 372
                               38.560 11.772 30.806 1.00 23.28
    ATOM 1246 CA LEU 372
15
                                38.517 11.709 32.343 1.00 27.76
    ATOM 1247 CB LEU 372
    ATOM 1248 CG LEU 372
                                37.155 11.306 32.924 1.00 21.18
                                37,289 10,891 34,381 1,00 27,64
    ATOM 1249 CD1 LEU 372
    ATOM 1250 CD2 LEU 372
                                36.197 12.480 32.763 1.00 20.90
                               39.804 12.505 30.357 1.00 21.34
    ATOM 1251 C LEU 372
20
                               39.779 13.708 30.086 1.00 23.16
    ATOM 1252 O LEU 372
                               40.896 11.761 30.276 1.00 24.42
    ATOM 1253 N LEU 373
                                42.177 12.302 29.855 1.00 23.78
    ATOM 1254 CA LEU 373
                                43,222 11.205 30.007 1.00 22.18
    ATOM 1255 CB LEU 373
    ATOM 1256 CG LEU 373
                                44.724 11.456 30.036 1.00 31.52
25
                                45.099 12.565 31.001 1.00 31.93
    ATOM 1257 CD1 LEU 373
                                45.382 10.152 30.460 1.00 30.24
    ATOM 1258 CD2 LEU 373
                               42.025 12.757 28.399 1.00 25.69
    ATOM 1259 C LEU 373
                               42.469 13.842 28.025 1.00 30.13
    ATOM 1260 O LEU 373
                               41.370 11.934 27.587 1.00 26.24
    ATOM 1261 N GLN 374
30
                                41.151 12.269 26.184 1.00 21.60
    ATOM 1262 CA GLN 374
                                40.501 11.091 25.444 1.00 24.57
    ATOM 1263 CB GLN 374
                                41.428 9.900 25.234 1.00 21.02
    ATOM 1264 CG GLN 374
                                40,762 8.744 24.501 1.00 22.86
    ATOM 1265 CD GLN 374
                                41.407 7.754 24.174 1.00 24.07
    ATOM 1266 OE1 GLN 374
35
                                39.466 8.865 24.249 1.00 25.59
    ATOM 1267 NE2 GLN 374
                               40,267 13,498 26,070 1,00 20,66
    ATOM 1268 C GLN 374
                               40.518 14.366 25.242 1.00 24.47
    ATOM 1269 O GLN 374
                               39.237 13.579 26.902 1.00 16.26
    ATOM 1270 N ALA 375
     ATOM 1271 CA ALA 375
                                38.337 14.727 26.870 1.00 17.16
40
                                37.156 14.491 27.803 1.00 19.53
    ATOM 1272 CB ALA 375
    ATOM 1273 C ALA 375
                               39.056 16.024 27.252 1.00 25.13
                               38.722 17.100 26.750 1.00 23.81
     ATOM 1274 O ALA 375
                               40.036 15.926 28.156 1.00 24.57
     ATOM 1275 N VAL 376
                                40.796 17.101 28.568 1.00 25.80
     ATOM 1276 CA VAL 376
45
                                41.711 16.792 29.814 1.00 26.48
     ATOM 1277 CB VAL 376
                                42.625 17.971 30.102 1.00 23.20
     ATOM 1278 CG1 VAL 376
     ATOM 1279 CG2 VAL 376
                                40.845 16.521 31.044 1.00 19.08
     ATOM 1280 C VAL 376
                               41.653 17.580 27.396 1.00 25.69
     ATOM 1281 O VAL 376
                               41.775 18.780 27.151 1.00 27.87
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WO 99/26966

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ATOM 1282 N LEU 377
                               42.249 16.637 26.666 1.00 23.09
     ATOM 1283 CA LEU 377
                                43.071 16.982 25.513 1.00 22.86
                                43.748 15.730 24.962 1.00 18.50
     ATOM 1284 CB LEU 377
                                44.814 15.096 25.867 1.00 22.65
     ATOM 1285 CG LEU 377
     ATOM 1286 CD1 LEU 377
                                45.144 13.708 25.374 1.00 16.70
 5
                                46.070 15.987 25.901 1.00 19.58
     ATOM 1287 CD2 LEU 377
                               42.197 17.634 24.430 1.00 26.14
     ATOM 1288 C LEU 377
     ATOM 1289 O LEU 377
                               42.579 18.638 23.830 1.00 20.62
     ATOM 1290 N LEU 378
                               41.016 17.057 24.208 1.00 28.99
     ATOM 1291 CA LEU 378
                                40.076 17.578 23.218 1.00 28.87
10
     ATOM 1292 CB LEU 378
                                38.814 16.710 23.182 1.00 26.89
                                37.637 17.167 22.311 1.00 28.83
     ATOM 1293 CG LEU 378
     ATOM 1294 CD1 LEU 378
                                38.053 17.273 20.840 1.00 27.97
     ATOM 1295 CD2 LEU 378
                                36.496 16.175 22.478 1.00 27.69
                               39.693 19.025 23.504 1.00 31.09
     ATOM 1296 C LEU 378
15
                               39.812 19.883 22.629 1.00 31.77
     ATOM 1297 O LEU 378
                                39.247 19.297 24.729 1.00 31.44
     ATOM 1298 N MET 379
     ATOM 1299 CA MET 379
                                38.841 20.649 25.104 1.00 32.62
                                37.876 20.603 26.293 1.00 31.45
     ATOM 1300 CB MET 379
     ATOM 1301 CG MET 379
                                36.586 19.855 26.010 1.00 38.75
20
     ATOM 1302 SD MET 379
                                35.646 20.541 24.601 1.00 41.27
     ATOM 1303 CE MET 379
                                34.231 19.443 24.609 1.00 35.68
                               39.980 21.613 25.421 1.00 33.72
     ATOM 1304 C MET 379
     ATOM 1305 O MET 379
                                39.940 22.297 26.446 1.00 36.29
     ATOM 1306 N SER 380
                               40.981 21.676 24.543 1.00 34.49
25
     ATOM 1307 CA SER 380
                                42.116 22.585 24.721 1.00 33.97
     ATOM 1308 CB SER 380
                                43.371 22.025 24.061 1.00 31.24
     ATOM 1309 OG SER 380
                                43.771 20.814 24.674 1.00 39.42
     ATOM 1310 C SER 380
                               41.772 23.926 24.088 1.00 39.69
                               41.787 24.069 22.864 1.00 44.64
     ATOM 1311 O SER 380
30
                               41.472 24.907 24.927 1.00 41.04
     ATOM 1312 N SER 381
                                41.090 26.234 24.462 1.00 44.91
     ATOM 1313 CA SER 381
                                40.406 27.004 25.594 1.00 44.50
     ATOM 1314 CB SER 381
                                41.294 27.177 26.678 1.00 45.42
     ATOM 1315 OG SER 381
     ATOM 1316 C SER 381
                               42.231 27.084 23.921 1.00 44.59
35
                               42.012 28.227 23.516 1.00 49.32
     ATOM 1317 O SER 381
                               43.440 26.541 23.896 1.00 43.75
     ATOM 1318 N ASP 382
                                44.571 27.315 23.407 1.00 43.93
     ATOM 1319 CA ASP 382
     ATOM 1320 CB ASP 382
                                45.817 27.047 24.257 1.00 48.39
     ATOM 1321 CG ASP 382
                                46.319 25.632 24.113 1.00 53.23
40
     ATOM 1322 OD1 ASP 382
                                45.590 24.702 24.517 1.00 56.97
     ATOM 1323 OD2 ASP 382
                                47.440 25.449 23.584 1.00 58.91
                               44.900 27.026 21.955 1.00 41.09
     ATOM 1324 C ASP 382
                               45.912 27.502 21.446 1.00 40.93
     ATOM 1325 O ASP 382
45
     ATOM 1326 N ARG 383
                                44.068 26.236 21.287 1.00 42.63
                                44.316 25.937 19.876 1.00 43.32
     ATOM 1327 CA ARG 383
                                43.289 24.935 19.331 1.00 42.31
     ATOM 1328 CB ARG 383
                                43.174 23.619 20.095 1.00 40.83
     ATOM 1329 CG ARG 383
     ATOM 1330 CD ARG 383
                                44.478 22.835 20.139 1.00 38.09
     ATOM 1331 NE ARG 383
                                44.271 21.542 20.787 1.00 37.33
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ATOM 1332 CZ ARG 383
                                45.235 20.690 21.115 1.00 38.35
                                 46.505 20.972 20.850 1.00 33.70
    ATOM 1333 NH1 ARG 383
                                 44.922 19.545 21.704 1.00 35.46
    ATOM 1334 NH2 ARG 383
                               44.166 27.256 19.127 1.00 44.96
    ATOM 1335 C ARG 383
    ATOM 1336 O ARG 383
                               43.214 28.006 19.361 1.00 45.60
                               45.112 27.574 18.230 1.00 45.33
    ATOM 1337 N PRO 384
                                46.330 26.852 17.836 1.00 46.85
    ATOM 1338 CD PRO 384
    ATOM 1339 CA PRO 384
                                45.024 28.830 17.484 1.00 47.37
                                46.323 28.823 16.672 1.00 46.90
    ATOM 1340 CB PRO 384
                                47.257 27.998 17.552 1.00 46.41
    ATOM 1341 CG PRO 384
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    ATOM 1342 C PRO 384
                               43.788 28.910 16.590 1.00 48.29
    ATOM 1343 O PRO 384
                               43.394 27.927 15.960 1.00 48.34
    ATOM 1344 N GLY 385
                               43.176 30.090 16.552 1.00 49.88
    ATOM 1345 CA GLY 385
                                42.013 30.290 15.712 1.00 50.35
                               40.669 29.958 16.324 1.00 50.70
    ATOM 1346 C GLY 385
15
                               39.639 30.201 15.697 1.00 53.48
    ATOM 1347 O GLY 385
                               40.663 29.404 17.529 1.00 49.04
    ATOM 1348 N LEU 386
                                39.405 29.057 18.182 1.00 50.53
    ATOM 1349 CA LEU 386
    ATOM 1350 CB LEU 386
                                39.655 28.433 19.558 1.00 45.17
    ATOM 1351 CG LEU 386
                                40.245 27.019 19.544 1.00 48.26
20
    ATOM 1352 CD1 LEU 386
                                40.502 26.564 20.970 1.00 41.68
                                39.285 26.065 18.836 1.00 38.40
    ATOM 1353 CD2 LEU 386
    ATOM 1354 C LEU 386
                               38.495 30.268 18.319 1.00 52.13
                               38.955 31.395 18.476 1.00 53.67
    ATOM 1355 O LEU 386
    ATOM 1356 N ALA 387
25
                               37.193 30.020 18.261 1.00 53.42
                                36.225 31.093 18.354 1.00 56.01
    ATOM 1357 CA ALA 387
    ATOM 1358 CB ALA 387
                                35.221 30.976 17.202 1.00 56.47
                               35.482 31.144 19.681 1.00 55.52
    ATOM 1359 C ALA 387
                               35.491 32.171 20.358 1.00 53.75
    ATOM 1360 O ALA 387
                               34.854 30.038 20.065 1.00 56.03
    ATOM 1361 N CYS 388
30
                                34.072 30.036 21.312 1.00 59.57
     ATOM 1362 CA CYS 388
    ATOM 1363 CB CYS 388
                                32.724 29.351 21.089 1.00 59.23
     ATOM 1364 SG CYS 388
                                31.314 30.363 21.641 1.00 58.64
                               34.846 29.289 22.398 1.00 62.18
     ATOM 1365 C CYS 388
                               34.458 28.190 22.790 1.00 67.88
     ATOM 1366 O CYS 388
35
                               35.955 29.950 22.760 1.00 60.78
    ATOM 1367 N VAL 389
                                37.005 29.583 23.713 1.00 57.70
    ATOM 1368 CA VAL 389
                                38.202 30.580 23.565 1.00 57.09
    ATOM 1369 CB VAL 389
    ATOM 1370 CG1 VAL 389
                                39.351 30.194 24.494 1.00 59.03
    ATOM 1371 CG2 VAL 389
                                38.671 30.618 22.124 1.00 53.98
40
                               36.661 29.515 25.195 1.00 57.77
     ATOM 1372 C VAL 389
     ATOM 1373 O VAL 389
                               36.943 28.513 25.851 1.00 60.94
    ATOM 1374 N GLU 390
                               36.102 30.594 25.732 1.00 52.68
     ATOM 1375 CA GLU 390
                                35.738 30.636 27.138 1.00 48.41
                                35.001 31.928 27.451 1.00 45.19
45
     ATOM 1376 CB GLU 390
                               34.868 29.439 27.459 1.00 47.63
     ATOM 1377 C GLU 390
     ATOM 1378 O GLU 390
                               34.986 28.837 28.529 1.00 51.95
                               34.002 29.082 26.517 1.00 47.11
     ATOM 1379 N ARG 391
                                33.099 27.950 26.699 1.00 51.64
     ATOM 1380 CA ARG 391
                                32.050 27.930 25.588 1.00 54.22
50
     ATOM 1381 CB ARG 391
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30.830 27.094 25.915 1.00 64.20
     ATOM 1382 CG ARG 391
     ATOM 1383 CD ARG 391
                                29.867 27.074 24.748 1.00 73.80
     ATOM 1384 NE ARG 391
                                28.533 26.622 25.128 1.00 79.76
     ATOM 1385 CZ ARG 391
                                27.714 27.298 25.929 1.00 84.27
    ATOM 1386 NH1 ARG 391
                                 28.090 28.465 26.439 1.00 85.28
     ATOM 1387 NH2 ARG 391
                                 26.515 26.809 26.217 1.00 86.84
                               33.890 26.644 26.684 1.00 48.18
     ATOM 1388 C ARG 391
                               33.504 25.671 27.330 1.00 49.57
     ATOM 1389 O ARG 391
                              34,987 26.625 25.936 1.00 45.01
     ATOM 1390 N ILE 392
                               35.835 25.440 25.858 1.00 48.77
     ATOM 1391 CA ILE 392
10
     ATOM 1392 CB ILE 392
                               36.854 25.565 24.692 1.00 46.45
                               37.798 24.370 24.679 1.00 42.35
     ATOM 1393 CG2 ILE 392
                               36.086 25.664 23.367 1.00 49.69
     ATOM 1394 CG1 ILE 392
                                36.950 25.897 22.136 1.00 51.09
     ATOM 1395 CD1 ILE 392
                              36.570 25.246 27.192 1.00 50.90
     ATOM 1396 C ILE 392
15
                              36,731 24.118 27.657 1.00 52.21
     ATOM 1397 O ILE 392
     ATOM 1398 N GLU 393
                               36.999 26.346 27.811 1.00 50.43
     ATOM 1399 CA GLU 393
                                37.673 26.267 29.101 1.00 50.30
     ATOM 1400 CB GLU 393
                                38.202 27.638 29.531 1.00 53.97
                                39.322 28.168 28.658 1.00 62.18
     ATOM 1401 CG GLU 393
20
                                39.911 29.478 29.168 1.00 67.69
     ATOM 1402 CD GLU 393
                                40.869 29.977 28.537 1.00 66.42
     ATOM 1403 OE1 GLU 393
     ATOM 1404 OE2 GLU 393
                                39.423 30.009 30.191 1.00 70.64
                               36.686 25.765 30.145 1.00 49.31
     ATOM 1405 C GLU 393
                               37.018 24.923 30.980 1.00 49.53
25
     ATOM 1406 O GLU 393
                               35.468 26.286 30.090 1.00 46.07
     ATOM 1407 N LYS 394
                                34.428 25.893 31.022 1.00 45.76
     ATOM 1408 CA LYS 394
                                33.147 26.666 30.727 1.00 43.85
     ATOM 1409 CB LYS 394
                               34.188 24.391 30.909 1.00 46.69
     ATOM 1410 C LYS 394
     ATOM 1411 O LYS 394
                               33.982 23.699 31.911 1.00 49.13
30
                               34.223 23.887 29.679 1.00 46.57
     ATOM 1412 N TYR 395
     ATOM 1413 CA TYR 395
                                34.014 22.467 29.427 1.00 43.33
     ATOM 1414 CB TYR 395.
                                33.818 22.211 27.929 1.00 48.44
     ATOM 1415 CG TYR 395
                                32.493 22.710 27.335 1.00 53.83
                                32.302 22.727 25.947 1.00 56.43
35
     ATOM 1416 CD1 TYR 395
                                31.078 23.148 25.374 1.00 59.73
     ATOM 1417 CE1 TYR 395
     ATOM 1418 CD2 TYR 395
                                 31.434 23.132 28.153 1.00 56.47
                                30,198 23.559 27.592 1.00 62.60
     ATOM 1419 CE2 TYR 395
     ATOM 1420 CZ TYR 395
                                30.037 23.562 26.200 1.00 63.18
     ATOM 1421 OH TYR 395
                                28.834 23.962 25.635 1.00 64.46
40
                               35.189 21.635 29.938 1.00 37.30
     ATOM 1422 C TYR 395
                               34.993 20.599 30.564 1.00 34.10
     ATOM 1423 O TYR 395
     ATOM 1424 N GLN 396
                               36.408 22.091 29.671 1.00 31.92
                                37.584 21.363 30.120 1.00 34.81
     ATOM 1425 CA GLN 396
                                38.861 21.987 29.560 1.00 32.64
     ATOM 1426 CB GLN 396
45
     ATOM 1427 CG GLN 396
                                40.114 21.183 29.882 1.00 29.57
                                41.370 21.827 29.352 1.00 29.46
     ATOM 1428 CD GLN 396
     ATOM 1429 OE1 GLN 396
                                41.648 22.982 29.649 1.00 34.65
     ATOM 1430 NE2 GLN 396
                                42.139 21.088 28.570 1.00 27.21
                               37.647 21.342 31.647 1.00 37.13
     ATOM 1431 C GLN 396
50
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	ATOM	1432 O GLN 396	37.939 20.302 32.236 1.00 37.36
	ATOM	1433 N ASP 397	37.371 22.484 32.284 1.00 38.61
	ATOM	1434 CA ASP 397	37.393 22.555 33.742 1.00 40.37
	ATOM	1435 CB ASP 397	37.099 23.973 34.240 1.00 40.51
5	ATOM	1436 CG ASP 397	38.130 24.974 33.772 1.00 43.77
,	ATOM	1437 OD1 ASP 397	39.330 24.632 33.775 1.00 46.50
	ATOM	1438 OD2 ASP 397	37.750 26.109 33.422 1.00 51.34
	ATOM	1439 C ASP 397	36.352 21.601 34.295 1.00 38.62
	ATOM	1440 O ASP 397	36.515 21.034 35.372 1.00 39.20
10	ATOM	1441 N SER 398	35.282 21.423 33.540 1.00 37.84
. 0	ATOM	1442 CA SER 398	34.221 20.524 33.942 1.00 37.80
	ATOM	1443 CB SER 398	33.039 20.669 32.984 1.00 34.28
	ATOM	1444 OG SER 398	31.981 19.815 33.360 1.00 46.60
	ATOM	1445 C SER 398	34.752 19.082 33.939 1.00 38.41
15	ATOM	1446 O SER 398	34.372 18.274 34.787 1.00 39.98
	ATOM	1447 N PHE 399	35.630 18.772 32.987 1.00 34.82
	ATOM	1448 CA PHE 399	36.213 17.433 32.885 1.00 35.96
	ATOM	1449 CB PHE 399	36.809 17.181 31.493 1.00 35.75
	ATOM	1450 CG PHE 399	35.775 16.936 30.419 1.00 39.30
20	<b>ATOM</b>	1451 CD1 PHE 399	35.640 17.826 29.344 1.00 39.86
	ATOM	1452 CD2 PHE 399	34.936 15.819 30.487 1.00 36.81
	ATOM	1453 CE1 PHE 399	34.674 17.607 28.330 1.00 41.25
	ATOM	1454 CE2 PHE 399	33.962 15.577 29.488 1.00 43.61
	ATOM	1455 CZ PHE 399	33.829 16.480 28.402 1.00 40.34
25	ATOM	1456 C PHE 399	37.306 17.217 33.921 1.00 33.48
	ATOM	1457 O PHE 399	37.406 16.139 34.512 1.00 26.86
	ATOM	1458 N LEU 400	38.132 18.239 34.118 1.00 31.47
	ATOM	1459 CA LEU 400	39.213 18.162 35.086 1.00 37.41
	ATOM	1460 CB LEU 400	40.051 19.441 35.038 1.00 34.24
30	ATOM	1461 CG LEU 400	40.934 19.574 33.788 1.00 35.10
	ATOM	1462 CD1 LEU 400	41.469 20.991 33.651 1.00 26.60
	ATOM	1463 CD2 LEU 400	42.077 18.569 33.884 1.00 29.44 38.666 17.931 36.491 1.00 38.84
	ATOM	1464 C LEU 400 1465 O LEU 400	
25	ATOM ATOM	1465 O LEU 400 1466 N LEU 401	39.137 17.049 37.205 1.00 40.38 37.654 18.703 36.870 1.00 42.79
35	ATOM	1467 CA LEU 401	37.056 18.584 38.197 1.00 43.48
	ATOM	1468 CB LEU 401	35.997 19.675 38.406 1.00 44.73
	ATOM	1469 CG LEU 401	35.322 19.737 39.779 1.00 51.39
	ATOM	1470 CD1 LEU 401	36.359 20.002 40.866 1.00 50.11
40	ATOM	1471 CD2 LEU 401	34.273 20.834 39.778 1.00 49.30
70	ATOM	1471 CD2 EEU 401	36.433 17.215 38.409 1.00 41.62
	ATOM	1473 O LEU 401	36.563 16.622 39.482 1.00 45.14
	ATOM	1474 N ALA 402	35.744 16.712 37.389 1.00 37.92
	ATOM		35.115 15.402 37.484 1.00 29.90
45	ATOM		34.196 15.187 36.297 1.00 30.70
	ATOM	1477 C ALA 402	36.203 14.336 37.508 1.00 28.88
	ATOM	1478 O ALA 402	36.083 13.322 38.188 1.00 32.14
	ATOM	1479 N PHE 403	37.274 14.588 36.764 1.00 31.07
	ATOM		38.402 13.656 36.661 1.00 29.90
50	ATOM	1481 CB PHE 403	39.396 14.178 35.605 1.00 27.03

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ATOM 1482 CG PHE 403
                                40.434 13.146 35.140 1.00 26.97
     ATOM 1483 CD1 PHE 403
                                41.362 13.509 34.149 1.00 25.55
     ATOM 1484 CD2 PHE 403
                                40.475 11.841 35.664 1.00 19.75
     ATOM 1485 CE1 PHE 403
                                42.331 12.588 33.679 1.00 27.90
     ATOM 1486 CE2 PHE 403
                                41.441 10.899 35.206 1.00 22.56
     ATOM 1487 CZ PHE 403
                                42.371 11.273 34.210 1.00 22.24
                               39.081 13.523 38.023 1.00 28.82
     ATOM 1488 C PHE 403
     ATOM 1489 O PHE 403
                               39.313 12.413 38.495 1.00 26.00
     ATOM 1490 N GLU 404
                               39.405 14.652 38.652 1.00 30.25
     ATOM 1491 CA GLU 404
                               40.039 14.627 39.966 1.00 34.03
10
     ATOM 1492 CB GLU 404
                                40.264 16.046 40.497 1.00 39.45
                                40.987 16.076 41.839 1.00 47.68
     ATOM 1493 CG GLU 404
     ATOM 1494 CD GLU 404
                                41.062 17.465 42.446 1.00 54.02
     ATOM 1495 OE1 GLU 404
                                41.607 18.380 41.796 1.00 57.27
                                40.573 17.638 43.585 1.00 63.85
15
     ATOM 1496 OE2 GLU 404
     ATOM 1497 C GLU 404
                               39.164 13.860 40.960 1.00 36.01
     ATOM 1498 O GLU 404
                               39.661 12.997 41.701 1.00 38.64
     ATOM 1499 N HIS 405
                              37.870 14.168 40.975 1.00 29.56
     ATOM 1500 CA HIS 405
                               36.949 13.508 41.892 1.00 31.69
20
     ATOM 1501 CB HIS 405
                               35.534 14.077 41.757 1.00 33.75
     ATOM 1502 CG HIS 405
                               35.401 15.498 42.213 1.00 34.75
     ATOM 1503 CD2 HIS 405
                                36.308 16.361 42.730 1.00 34.58
     ATOM 1504 ND1 HIS 405
                                34.207 16.187 42.146 1.00 32.43
     ATOM 1505 CE1 HIS 405
                                34.385 17.414 42.598 1.00 36.15
25
     ATOM 1506 NE2 HIS 405
                                35.650 17.549 42.960 1.00 39.84
     ATOM 1507 C HIS 405
                              36.904 12.013 41.673 1.00 34.21
     ATOM 1508 O HIS 405
                              36.700 11.247 42.624 1.00 37.06
     ATOM 1509 N TYR 406
                               37.081 11.594 40.419 1.00 30.83
     ATOM 1510 CA TYR 406
                                37.059 10.173 40.093 1.00 28.85
     ATOM 1511 CB TYR 406
                                37.018 9.959 38.575 1.00 31.48
30
     ATOM 1512 CG TYR 406
                                36.879 8.490 38.181 1.00 23.49
                                35.683 7.798 38.397 1.00 19.42
     ATOM 1513 CD1°TYR 406
     ATOM 1514 CE1 TYR 406
                                35.556 6.427 38.059 1.00 23.80
     ATOM 1515 CD2 TYR 406
                                37.950 7.794 37.624 1.00 21.81
35
    ATOM 1516 CE2 TYR 406
                                37.838 6.421 37.278 1.00 24.64
    ATOM 1517 CZ TYR 406
                                36.639 5.753 37.503 1.00 21.56
    ATOM 1518 OH TYR 406
                                36.537 4.404 37.186 1.00 24.96
    ATOM 1519 C TYR 406
                               38.318 9.526 40.638 1.00 24.24
    ATOM 1520 O TYR 406
                               38.308 8.375 41.050 1.00 27.08
40
    ATOM 1521 N ILE 407
                              39.407 10.278 40.617 1.00 25.76
    ATOM 1522 CA ILE 407
                               40.688 9.799 41.105 1.00 33.75
    ATOM 1523 CB ILE 407
                               41.815 10.822 40.796 1.00 34.23
    ATOM 1524 CG2 ILE 407
                               43.121 10.400 41.435 1.00 32.46
    ATOM 1525 CG1 ILE 407
                               41.959 10.972 39.275 1.00 43.30
                               42.267 9.677 38.523 1.00 40.40
45
    ATOM 1526 CD1 ILE 407
    ATOM 1527 C ILE 407
                              40.620 9.556 42.613 1.00 39.03
    ATOM 1528 O ILE 407
                              41.192 8.583 43.107 1.00 35.18
    ATOM 1529 N ASN 408
                               39.916 10.440 43.335 1.00 37.25
    ATOM 1530 CA ASN 408
                               39.778 10.292 44.777 1.00 37.01
50
    ATOM 1531 CB ASN 408
                               39.099 11.514 45.400 1.00 32.27
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	ATOM	1532 CG ASN 408 39.887 12.790 45.187 1.00 33.56
	ATOM	
	ATOM	1534 ND2 ASN 408 39.182 13.903 44.996 1.00 31.23
	ATOM	1535 C ASN 408 38.961 9.046 45.055 1.00 38.14
5	ATOM	1536 O ASN 408 39,303 8.243 45,920 1.00 42.16
	ATOM	1537 N TYR 409 37.874 8.894 44.303 1.00 35.62
	ATOM	1538 CA TYR 409 37,002 7.733 44.412 1.00 35.91
	ATOM	1539 CB TYR 409 35.929 7.804 43.323 1.00 34.41
	ATOM	1540 CG TYR 409 35.196 6.495 43.066 1.00 38.73
10	ATOM	1541 CD1 TYR 409 34.266 5.982 43.980 1.00 41.34
	ATOM	1542 CE1 TYR 409 33.600 4.745 43.741 1.00 47.16
	ATOM	1543 CD2 TYR 409 35.461 5.752 41.907 1.00 46.20
	ATOM	1544 CE2 TYR 409 34.814 4.518 41.651 1.00 50.74
	ATOM	1545 CZ TYR 409 33.891 4.023 42.573 1.00 50.88
15	ATOM	1546 OH TYR 409 33.262 2.816 42.302 1.00 53.14
_	ATOM	1547 C TYR 409 37.827 6.459 44.240 1.00 38.16
	ATOM	1548 O TYR 409 37.806 5.561 45.082 1.00 41.83
	ATOM	1549 N ARG 410 38.551 6.399 43.125 1.00 42.25
	ATOM	1550 CA ARG 410 39.410 5.272 42.765 1.00 42.83
20	ATOM	1551 CB ARG 410 40.029 5.540 41.392 1.00 36.83
	ATOM	1552 CG ARG 410 39.055 5.397 40.249 1.00 34.32
	ATOM	1553 CD ARG 410 39.134 3.996 39.681 1.00 36.62
	<b>ATOM</b>	1554 NE ARG 410 40.420 3.787 39.013 1.00 38.64
	<b>ATOM</b>	1555 CZ ARG 410 40.832 2.625 38.517 1.00 35.73
25	ATOM	1556 NH1 ARG 410 40.068 1.548 38.617 1.00 33.17
	<b>ATOM</b>	1557 NH2 ARG 410 42.006 2.544 37.916 1.00 32.70
	<b>ATOM</b>	1558 C ARG 410 40.520 5.039 43.780 1.00 46.67
	ATOM	1559 O ARG 410 40.900 3.901 44.053 1.00 41.78
	<b>ATOM</b>	1560 N LYS 411 41.026 6.140 44.325 1.00 52.99
30	ATOM	1561 CA LYS 411 42.109 6.141 45.298 1.00 58.32
	ATOM	1562 CB LYS 411 41.565 5.956 46.731 1.00 64.99
	ATOM	1563 CG LYS 411 40.660 4.763 46.977 1.00 70.48
	ATOM	1564 CD LYS 411 40.034 4.866 48.364 1.00 77.18
	ATOM	1565 CE LYS 411 39.053 3.732 48.625 1.00 84.30
35	ATOM	1566 NZ LYS 411 38.392 3.865 49.958 1.00 86.48
	ATOM	1567 C LYS 411 43.238 5.163 45.000 1.00 56.66
	ATOM	1568 O LYS 411 43.329 4.075 45.575 1.00 55.47
	ATOM	1569 N HIS 412 44.091 5.582 44.070 1.00 54.67
	ATOM	1570 CA HIS 412 45.266 4.823 43.657 1.00 48.67
40	<b>ATOM</b>	1571 CB HIS 412 45.878 5.442 42.393 1.00 43.14
		1572 CG HIS 412 45.073 5.218 41.156 1.00 41.36
	<b>ATOM</b>	1573 CD2 HIS 412 44.084 5.952 40.584 1.00 35.44
	<b>ATOM</b>	1574 ND1 HIS 412 45.220 4.093 40.364 1.00 38.19
	<b>ATOM</b>	1575 CE1 HIS 412 44.357 4.150 39.363 1.00 34.75
45	<b>ATOM</b>	1576 NE2 HIS 412 43.659 5.263 39.474 1.00 35.52
	<b>ATOM</b>	1577 C HIS 412 46.264 4.932 44.793 1.00 46.35
	<b>ATOM</b>	1578 O HIS 412 46.326 5.951 45.479 1.00 42.73
	ATOM	1579 N HIS 413 47.049 3.883 44.993 1.00 48.92
		1580 CA HIS 413 48.040 3.903 46.052 1.00 53.15
50		1581 CB HIS 413 48.148 2.515 46.688 1.00 55.27

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ATOM 1582 CG HIS 413
                                46.843 2.015 47.238 1.00 58.77
     ATOM 1583 CD2 HIS 413
                                46.138 0.892 46.977 1.00 61.65
     ATOM 1584 ND1 HIS 413
                                46.108 2.726 48.161 1.00 60.31
                                45.003 2.061 48.445 1.00 63.01
     ATOM 1585 CE1 HIS 413
                                44.993 0.942 47.743 1.00 62.93
     ATOM 1586 NE2 HIS 413
                               49.359 4.364 45.456 1.00 53.19
     ATOM 1587 C HIS 413
                               50.335 3.617 45.390 1.00 54.93
     ATOM 1588 O HIS 413
     ATOM 1589 N VAL 414
                                49.343 5.612 44.999 1.00 53.77
     ATOM 1590 CA VAL 414
                                50.487 6.282 44.389 1.00 51.06
     ATOM 1591 CB VAL 414
                                50.374 6.305 42.838 1.00 51.49
10
     ATOM 1592 CG1 VAL 414
                                 51.603 6.958 42.231 1.00 45.22
     ATOM 1593 CG2 VAL 414
                                 50.210 4.891 42.304 1.00 52.67
     ATOM 1594 C VAL 414
                               50.444 7.724 44.894 1.00 54.28
     ATOM 1595 O VAL 414
                                49.418 8.401 44.774 1.00 55.49
15
     ATOM 1596 N THR 415
                               51.547 8.190 45.467 1.00 56.28
     ATOM 1597 CA THR 415
                                51.610 9.550 45.986 1.00 57.83
     ATOM 1598 CB THR 415
                                52.874 9.756 46.858 1.00 59.64
     ATOM 1599 OG1 THR 415
                                 52.922 11.115 47.311 1.00 66.69
     ATOM 1600 CG2 THR 415
                                 54.137 9.436 46.067 1.00 59.42
20
     ATOM 1601 C THR 415
                               51.599 10.577 44.855 1.00 56.98
     ATOM 1602 O THR 415
                               52.176 10.345 43.789 1.00 55.70
     ATOM 1603 N HIS 416
                               50.936 11.707 45.093 1.00 57.44
     ATOM 1604 CA HIS 416
                               50.835 12.786 44.108 1.00 57.34
     ATOM 1605 CB HIS 416
                               52.207 13.425 43.875 1.00 61.35
25
     ATOM 1606 CG HIS 416
                               52.860 13.940 45.123 1.00 69.78
     ATOM 1607 CD2 HIS 416
                                54.049 13.633 45.695 1.00 71.42
     ATOM 1608 ND1 HIS 416
                                52.283 14.901 45.922 1.00 72.49
     ATOM 1609 CE1 HIS 416
                                53.087 15.165 46.938 1.00 75.50
     ATOM 1610 NE2 HIS 416
                                54.165 14.410 46.819 1.00 73.91
     ATOM 1611 C HIS 416
30
                              50.301 12.260 42.773 1.00 53.79
     ATOM 1612 O HIS 416
                               50.769 12.667 41.710 1.00 52.81
     ATOM 1613 N PHE 417
                               49.318 11.366 42.824 1.00 48.05
     ATOM 1614 CA PHE 417
                                48.769 10.784 41.610 1.00 47.99
     ATOM 1615 CB PHE 417
                                47.652 9.799 41.940 1.00 46.11
35
     ATOM 1616 CG PHE 417
                                47.314 8.868 40.791 1.00 44.27
     ATOM 1617 CD1 PHE 417
                                48.155 7.796 40.481 1.00 41.79
     ATOM 1618 CD2 PHE 417
                                46.179 9.091 40.003 1.00 40.23
     ATOM 1619 CE1 PHE 417
                                47.872 6.936 39.386 1.00 44.30
                                45.874 8.248 38.907 1.00 36.80
     ATOM 1620 CE2 PHE 417
40
     ATOM 1621 CZ PHE 417
                                46.725 7.167 38.595 1.00 40.69
    ATOM 1622 C PHE 417
                               48.227 11.824 40.625 1.00 46.69
    ATOM 1623 O PHE 417
                               48.551 11.787 39.436 1.00 43.35
    ATOM 1624 N TRP 418
                               47,410 12,746 41,124 1,00 45,14
    ATOM 1625 CA TRP 418 · 46.821 13.775 40.276 1.00 44.89
45
    ATOM 1626 CB TRP 418
                                45.808 14.604 41.077 1.00 42.24
    ATOM 1627 CG TRP 418
                                45.096 15.646 40.259 1.00 47.11
    ATOM 1628 CD2 TRP 418
                                44.186 15.417 39.159 1.00 46.98
    ATOM 1629 CE2 TRP 418
                                43.786 16.678 38.676 1.00 48.94
    ATOM 1630 CE3 TRP 418
                                43.676 14.261 38.548 1.00 45.23
50
    ATOM 1631 CD1 TRP 418
                                45.204 17.003 40.387 1.00 46.24
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	ATOM	1632 NE1 TRP 418	44.425 17.637 39.448 1.00 50.63
	ATOM	1633 CZ2 TRP 418	42.891 16.839 37.598 1.00 45.46
	ATOM	1634 CZ3 TRP 418	42.780 14.411 37.468 1.00 44.50
	ATOM	1635 CH2 TRP 418	42.403 15.696 37.009 1.00 47.55
5	ATOM	1636 C TRP 418	47.862 14.676 39.598 1.00 43.88
	ATOM	1637 O TRP 418	47.834 14.842 38.383 1.00 43.17
	ATOM	1638 N PRO 419	48.788 15.281 40.369 1.00 43.55
	ATOM	1639 CD PRO 419	49.006 15.290 41.826 1.00 41.52
	ATOM	1640 CA PRO 419	49.787 16.135 39.725 1.00 41.48
10	ATOM	1641 CB PRO 419	50.626 16.627 40.912 1.00 39.21
	ATOM	1642 CG PRO 419	49.593 16.667 42.017 1.00 39.25
	ATOM	1643 C PRO 419	50.616 15.363 38.701 1.00 36.28
	ATOM	1644 O PRO 419	50.940 15.882 37.638 1.00 37.08
	ATOM	1645 N LYS 420	50.959 14.124 39.033 1.00 35.96
15	ATOM	1646 CA LYS 420	51.742 13.281 38.132 1.00 40.82
	ATOM	1647 CB LYS 420	52.094 11.945 38.792 1.00 40.78
	ATOM	1648 CG LYS 420	53.086 12.046 39.933 1.00 48.62
	ATOM	1649 CD LYS 420	53.391 10.668 40.497 1.00 55.12
	ATOM	1650 CE LYS 420	54.395 10.741 41.635 1.00 53.26
20	ATOM	1651 NZ LYS 420	54.719 9.388 42.152 1.00 52.69
	ATOM	1652 C LYS 420	50.957 13.005 36.860 1.00 40.29
	ATOM	1653 O LYS 420	51.516 12.989 35.764 1.00 39.66
	ATOM	1654 N LEU 421	49.658 12.786 37.023 1.00 38.33
	ATOM	1655 CA LEU 421	48.784 12.507 35.903 1.00 37.60
25	ATOM	1656 CB LEU 421	47.417 12.074 36.428 1.00 43.66
	ATOM	1657 CG LEU 421	46.386 11.479 35.474 1.00 46.50
	ATOM	1658 CD1 LEU 421	46.946 10.253 34.770 1.00 45.15
	ATOM	1659 CD2 LEU 421	45.154 11.107 36.279 1.00 51.31
	ATOM	1660 C LEU 421	48.661 13.747 35.014 1.00 39.59
30	ATOM	1661 O LEU 421	48.599 13.638 33.791 1.00 40.66
	ATOM	1662 N LEU 422	48.642 14.928 35.623 1.00 39.57
	ATOM	1663 CA LEU 422	48.545 16.170 34.867 1.00 38.63
	ATOM	1664 CB LEU 422	48.313 17.357 35.802 1.00 41.79
	ATOM	1665 CG LEU 422	46.996 17.407 36.581 1.00 42.74
35	ATOM	1666 CD1 LEU 422	47.010 18.606 37.515 1.00 42.89
	ATOM	1667 CD2 LEU 422	45.823 17.494 35.628 1.00 39.27
	ATOM	1668 C LEU 422	49.808 16.410 34.039 1.00 40.47
	ATOM	1669 O LEU 422	49.747 17.029 32.979 1.00 47.83
	ATOM	1670 N MET 423	50.949 15.936 34.519 1.00 34.27
40	ATOM	1671 CA MET 423	52.187 16.103 33.774 1.00 35.25
	ATOM	1672 CB MET 423	53.403 15.716 34.622 1.00 32.56
	ATOM	1673 CG MET 423	53.675 16.654 35.774 1.00 40.70
	ATOM	1674 SD MET 423	55.226 16.278 36.597 1.00 47.65
	ATOM	1675 CE MET 423	54.920 14.601 37.163 1.00 47.16
45	ATOM	1676 C MET 423	52.164 15.254 32.502 1.00 35.13
	ATOM	1677 O MET 423	52.934 15.499 31.570 1.00 29.85
	ATOM	1678 N LYS 424	51.285 14.252 32.482 1.00 31.56
	ATOM	1679 CA LYS 424	51.152 13.384 31.316 1.00 32.29
	ATOM	1680 CB LYS 424	50.373 12.115 31.681 1.00 30.56
50	ATOM	1681 CG LYS 424	51.106 11.178 32.631 1.00 30.07

	ATOM	1682 CD LYS 424	52.248 10.482 31.938 1.00 33.22
	ATOM	1683 CE LYS 424	53.059 9.593 32.875 1.00 28.75
	ATOM	1684 NZ LYS 424	53.868 10.383 33.833 1.00 31.01
	ATOM	1685 C LYS 424	50.435 14.150 30.197 1.00 29.26
5	ATOM	1686 O LYS 424	50.719 13.944 29.030 1.00 30.22
_	ATOM	1687 N VAL 425	49.514 15.036 30.573 1.00 23.53
	ATOM	1688 CA VAL 425	48.792 15.849 29.601 1.00 28.91
	ATOM	1689 CB VAL 425	47.808 16.829 30.295 1.00 29.44
	ATOM	1690 CG1 VAL 425	47.148 17.737 29.273 1.00 28.81
10	ATOM	1691 CG2 VAL 425	46.744 16.049 31.057 1.00 31.22
. •	ATOM	1692 C VAL 425	49.822 16.669 28.831 1.00 32.03
	ATOM	1693 O VAL 425	49.771 16.769 27.605 1.00 31.95
	ATOM	1694 N THR 426	50,763 17.247 29.570 1.00 33.61
	ATOM	1695 CA THR 426	51.821 18.057 28.995 1.00 30.76
15	ATOM	1696 CB THR 426	52.678 18.695 30.105 1.00 32.34
	ATOM	1697 OG1 THR 426	51,842 19,535 30,912 1.00 33.07
	ATOM	1698 CG2 THR 426	53.812 19.533 29.514 1.00 25.40
	ATOM	1699 C THR 426	52.712 17.225 28.086 1.00 32.53
	ATOM	1700 O THR 426	53.113 17.686 27.014 1.00 35.19
20	ATOM	1701 N ASP 427	53.022 16.003 28.507 1.00 28.83
	ATOM	1702 CA ASP 427	53.858 15.130 27.695 1.00 35.12
	ATOM	1703 CB ASP 427	54.273 13.880 28.476 1.00 39.14
	ATOM	1704 CG ASP 427	55.122 14.212 29.693 1.00 45.80
	ATOM	1705 OD1 ASP 427	56.052 15.034 29.556 1.00 41.97
25	<b>ATOM</b>	1706 OD2 ASP 427	54.869 13.642 30.775 1.00 50.06
	<b>ATOM</b>	1707 C ASP 427	53.124 14.726 26.422 1.00 33.94
	<b>ATOM</b>	1708 O ASP 427	53.737 14.617 25.362 1.00 38.02
	<b>ATOM</b>	1709 N LEU 428	51.818 14.512 26.529 1.00 27.15
	<b>ATOM</b>	1710 CA LEU 428	51.013 14.148 25.373 1.00 29.99
30	<b>ATOM</b>	1711 CB LEU 428	49.602 13.719 25.802 1.00 22.49
	<b>ATOM</b>	1712 CG LEU 428	49.541 12.285 26.359 1.00 25.54
	<b>ATOM</b>	1713 CD1 LEU 428	48.210 12.021 27.037 1.00 20.60
	<b>ATOM</b>	1714 CD2 LEU 428	49.785 11.303 25.224 1.00 17.24
	ATOM	1715 C LEU 428	50.947 15.305 24.381 1.00 28.94
35	ATOM	1716 O LEU 428	50.941 15.088 23.174 1.00 31.26
	ATOM	1717 N ARG 429	50.910 16.531 24.887 1.00 27.64
	ATOM	1718 CA ARG 429	50.877 17.694 24.011 1.00 28.13
	ATOM	1719 CB ARG 429	50.584 18.969 24.800 1.00 29.59
	ATOM	1720 CG ARG 429	49.224 18.980 25.455 1.00 34.85
40	ATOM	1721 CD ARG 429	48.951 20.314 26.118 1.00 47.18
	ATOM	1722 NE ARG 429	47.657 20.358 26.797 1.00 57.93
	ATOM	1723 CZ ARG 429	46.473 20.193 26.200 1.00 63.62
	ATOM	1724 NH1 ARG 429	46.402 19.972 24.889 1.00 60.71
	ATOM	1725 NH2 ARG 429	45.356 20.257 26.919 1.00 62.38
45	ATOM	1726 C ARG 429	52.229 17.819 23.304 1.00 29.81
	ATOM	1727 O ARG 429	52.294 18.209 22.143 1.00 30.81
	ATOM	1728 N MET 430	53.305 17.482 24.008 1.00 29.64
	ATOM	1729 CA MET 430	54.639 17.545 23.422 1.00 34.72
	ATOM	1730 CB MET 430	55.716 17.323 24.485 1.00 34.97
50	ATOM	1731 CG MET 430	55.864 18.480 25.451 1.00 45.34

## . WO 99/26966

	ATOM	1732 SD MET 430	56.162 20.050 24.596 1.00 52.55
	<b>ATOM</b>	1733 CE MET 430	57.598 19.639 23.589 1.00 55.56
	<b>ATOM</b>	1734 C MET 430	54.778 16.500 22.325 1.00 34.01
	<b>ATOM</b>	1735 O MET 430	55.440 16.733 21.318 1.00 37.29
5	ATOM	1736 N ILE 431	54.161 15.340 22.533 1.00 29.99
	ATOM	1737 CA ILE 431	54.197 14.279 21.545 1.00 28.82
	<b>ATOM</b>	1738 CB ILE 431	53.523 12.984 22.095 1.00 27.39
	<b>ATOM</b>	1739 CG2 ILE 431	53.260 11.989 20.956 1.00 23.87
	<b>ATOM</b>	1740 CG1 ILE 431	54.414 12.386 23.201 1.00 25.56
10	ATOM	1741 CD1 ILE 431	53.850 11.155 23.896 1.00 17.29
	<b>ATOM</b>	1742 C ILE 431	53.450 14.785 20.301 1.00 29.49
	ATOM	1743 O ILE 431	53.908 14.603 19.174 1.00 24.19
	<b>ATOM</b>	1744 N GLY 432	52.311 15.435 20.524 1.00 25.25
	<b>ATOM</b>	1745 CA GLY 432	51.542 15.971 19.419 1.00 30.38
15	ATOM	1746 C GLY 432	52.334 16.997 18.614 1.00 32.75
	ATOM	1747 O GLY 432	52,410 16.895 17.387 1.00 36.38
	ATOM	1748 N ALA 433	52.930 17.974 19.294 1.00 26.77
	ATOM	1749 CA ALA 433	53.711 19.012 18.625 1.00 26.48
	ATOM	1750 CB ALA 433	54.182 20.047 19.631 1.00 19.90
20	ATOM	1751 C ALA 433	54.902 18.407 17.890 1.00 30.73
	ATOM	1752 O ALA 433	55.207 18.787 16.760 1.00 31.60
	ATOM	1753 N CYS 434	55.582 17.467 18.537 1.00 33.22
	ATOM	1754 CA CYS 434	56.728 16.801 17.914 1.00 34.34
	<b>ATOM</b>	1755 CB CYS 434	57.339 15.808 18.895 1.00 35.20
25	ATOM	1756 SG CYS 434	59.191 15.745 18.798 1.00 54.48
	ATOM	1757 C CYS 434	56.313 16.052 16.636 1.00 34.09
	<b>ATOM</b>	1758 O CYS 434	57.095 15.937 15.679 1.00 34.89
	<b>ATOM</b>	1759 N HIS 435	55.088 15.545 16.642 1.00 34.30
	ATOM	1760 CA HIS 435	54.570 14.818 15.501 1.00 35.44
30	ATOM	1761 CB HIS 435	53.296 14.061 15.886 1.00 31.76
	ATOM	1762 CG HIS 435	52.587 13.469 14.715 1.00 32.03
	ATOM	1763 CD2 HIS 435	52.735 12.277 14.092 1.00 28.61
	<b>ATOM</b>	1764 ND1 HIS 435	51.665 14.177 13.970 1.00 28.48
	ATOM	1765 CE1 HIS 435	51.284 13.453 12.941 1.00 33.27
35	<b>ATOM</b>	1766 NE2 HIS 435	51.920 12.284 12.985 1.00 31.57
	ATOM	1767 C HIS 435	54.311 15.750 14.319 1.00 32.74
	ATOM	1768 O HIS 435	54.504 15.363 13.175 1.00 32.87
	ATOM	1769 N ALA 436	53.881 16.975 14.608 1.00 31.01
	ATOM	1770 CA ALA 436	53.628 17.966 13.571 1.00 29.91
40	ATOM	1771 CB ALA 436	53.221 19.290 14.197 1.00 21.23
	<b>ATOM</b>	1772 C ALA 436	
	ATOM	1773 O ALA 436	
	<b>ATOM</b>	1774 N SER 437	56.030 18.266 13.483 1.00 35.19
	ATOM	1775 CA SER 437	57.344 18.426 12.871 1.00 33.03
45	ATOM	1776 CB SER 437	58.389 18.720 13.941 1.00 35.31
	ATOM	1777 OG SER 437	59.681 18.782 13.373 1.00 44.99
	ATOM	1778 C SER 437	57.758 17.178 12.100 1.00 38.39
	ATOM	1779 O SER 437	58.374 17.269 11.034 1.00 37.54
	ATOM	1780 N ARG 438	
50	ATOM	1781 CA ARG 438	57.762 14.754 11.992 1.00 39.30

# . WO 99/26966

	ATOM	1782 CB ARG 438	57.517 13.572 12.941 1.00 42.97
	ATOM	1783 CG ARG 438	58.542 13.436 14.059 1.00 41.72
	ATOM	1784 CD ARG 438	59.926 13.212 13.484 1.00 45.23
	ATOM	1785 NE ARG 438	59.961 12.050 12.601 1.00 45.66
5	ATOM	1786 CZ ARG 438	60.935 11.804 11.731 1.00 49.71
_	ATOM	1787 NH1 ARG 438	61.961 12.641 11.627 1.00 50.91
	ATOM	1788 NH2 ARG 438	60.885 10.727 10.960 1.00 46.86
	ATOM		56.939 14.565 10.725 1.00 42.37
	ATOM	1790 O ARG 438	57.311 13.794 9.841 1.00 40.58
10	ATOM		55.816 15.269 10.645 1.00 42.25
10	ATOM	1792 CA PHE 439	54.957 15.170 9.479 1.00 42.81
	ATOM	1793 CB PHE 439	53.593 15.790 9.771 1.00 42.18
	ATOM	1794 CG PHE 439	52.594 15.597 8.656 1.00 42.48
	ATOM	1795 CD1 PHE 439	52.173 14.312 8.295 1.00 47.09
15	ATOM	1796 CD2 PHE 439	52.086 16.696 7.961 1.00 39.76
13	ATOM	1797 CEI PHE 439	51.256 14.110 7.234 1.00 49.17
	ATOM	1797 CEI FHE 439	51.174 16.524 6.896 1.00 45.10
	ATOM	1798 CE2 FHE 439	50.751 15.225 6.532 1.00 46.36
	ATOM		55.626 15.905 8.322 1.00 44.79
20	ATOM		55.596 15.444 7.181 1.00 40.26
20	ATOM		56.236 17.049 8.629 1.00 42.77
	ATOM	1803 CA LEU 440	56.927 17.839 7.621 1.00 42.96
	ATOM	1804 CB LEU 440	57.421 19.156 8.216 1.00 37.19
	ATOM	1805 CG LEU 440	56.348 20.117 8.725 1.00 36.97
25	ATOM	1806 CD1 LEU 440	57.020 21.338 9.321 1.00 33.65
23	ATOM	1807 CD2 LEU 440	55.411 20.519 7.572 1.00 35.42
	ATOM		58.106 17.063 7.053 1.00 45.47
	ATOM		58.421 17.191 5.876 1.00 52.48
	ATOM		58.760 16.266 7.890 1.00 49.15
30	ATOM	1811 CA HIS 441	59.893 15.473 7.435 1.00 54.76
30			60.723 14.964 8.624 1.00 56.68
	ATOM	1812 CB HIS 441	61.515 16.026 9.323 1.00 62.73
	ATOM	1814 CD2 HIS 441	62.851 16.166 9.508 1.00 65.73
	ATOM	1815 ND1 HIS 441	60.929 17.098 9.966 1.00 66.01
25	ATOM	1816 CE1 HIS 441	61.871 17.845 10.518 1.00 65.55
35	ATOM ATOM	1817 NE2 HIS 441	63.044 17.306 10.258 1.00 60.09
			69.417 14.292 6.589 1.00 55.93
	ATOM ATOM		50.084 13.908 5.630 1.00 57.33
	ATOM		58.271 13.716 6.948 1.00 57.81
40			57.712 12.585 6.203 1.00 59.11
40	ATOM	1821 CA MET 442	
	ATOM	1822 CB MET 442	
	ATOM	1823 CG MET 442	
	ATOM	1824 SD MET 442	* * ** *
4 =	ATOM	1825 CE MET 442	
45	ATOM	1826 C MET 442	57.178 13.065 4.854 1.00 60.31 57,279 12.369 3.846 1.00 58.18
	ATOM	1827 O MET 442	
	ATOM		
	ATOM	1829 CA LYS 443	
50	ATOM	1830 CB LYS 443	
50	ATOM	1831 CG LYS 443	54.589 16.872 2.945 1.00 69.12

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54.064 18.250 3.363 1.00 71.14
    ATOM 1832 CD LYS 443
                               53.138 18.183 4.575 1.00 73.43
    ATOM 1833 CE LYS 443
                               52.668 19.534 5.015 1.00 67.97
    ATOM 1834 NZ LYS 443
                              57.112 15.030 2.585 1.00 67.29
    ATOM 1835 C LYS 443
                              56.800 15.218 1.406 1.00 67.90
    ATOM 1836 O LYS 443
    ATOM 1837 N VAL 444
                               58.373 14.941 2.996 1.00 66.57
    ATOM 1838 CA VAL 444
                               59.501 15.064 2.078 1.00 64.76
                               60.618 15.940 2.693 1.00 62.76
    ATOM 1839 CB VAL 444
                                61.767 16.092 1.712 1.00 64.00
    ATOM 1840 CG1 VAL 444
                                60.062 17.301 3.072 1.00 59.27
10
    ATOM 1841 CG2 VAL 444
    ATOM 1842 C VAL 444
                               60.091 13.693 1.744 1.00 68.61
    ATOM 1843 O VAL 444
                               60.145 13.294 0.577 1.00 70.60
                               60.520 12.972 2.775 1.00 70.71
    ATOM 1844 N GLU 445
    ATOM 1845 CA GLU 445
                               61.129 11.653 2.609 1.00 71.45
                               61.808 11.233 3.916 1.00 72.36
    ATOM 1846 CB GLU 445
15
    ATOM 1847 C GLU 445
                               60.181 10.547 2.148 1.00 71.46
    ATOM 1848 O GLU 445
                               60.588 9.390 2.042 1.00 73.02
    ATOM 1849 N CYS 446
                               58.925 10.895 1.871 1.00 71.12
    ATOM 1850 CA CYS 446
                               57.945 9.901 1.419 1.00 70.83
                               57.031 9.485 2.581 1.00 71.05
    ATOM 1851 CB CYS 446
20
    ATOM 1852 SG CYS 446
                               57.845 8.593 3.925 1.00 72.83
                              57.081 10.390 0.261 1.00 71.91
    ATOM 1853 C CYS 446
                               56.776 11.582 0.155 1.00 72.06
    ATOM 1854 O CYS 446
    ATOM 1855 N PRO 447
                               56.673 9.470 -0.635 1.00 73.12
    ATOM 1856 CD PRO 447
                               56.967 8.026 -0.671 1.00 72.88
25
    ATOM 1857 CA PRO 447
                               55.837 9.825 -1.784 1.00 74.22
                               55.717 8.500 -2.537 1.00 72.98
    ATOM 1858 CB PRO 447
    ATOM 1859 CG PRO 447
                              -57.015 7.790 -2.161 1.00 74.77
    ATOM 1860 C PRO 447
                               54.479 10.343 -1.330 1.00 75.94
                               53.754 9.652 -0.616 1.00 76.67
    ATOM 1861 O PRO 447
30
                               54.145 11.558 -1.755 1.00 76.91
    ATOM 1862 N THR 448
                              52.879 12.197 -1.403 1.00 78.24
    ATOM 1863 CA THR 448
                               52.647 13.459 -2.261 1.00 81.33
    ATOM 1864 CB THR 448
    ATOM 1865 OG1 THR 448
                               52.552 13.087 -3.643 1.00 84.46
                                53.802 14.444 -2.089 1.00 83.51
    ATOM 1866 CG2 THR 448
35
    ATOM 1867 C THR 448
                               51.676 11.270 -1.580 1.00 77.42
    ATOM 1868 O THR 448
                               50.662 11.413 -0.894 1.00 77.65
    ATOM 1869 N GLU 449
                               51.795 10.319 -2.502 1.00 76.29
    ATOM 1870 CA GLU 449
                               50.720 9.375 -2.783 1.00 75.03
                               51.048 8.572 -4.043 1.00 74.62
    ATOM 1871 CB GLU 449
40
                               50.445 8.421 -1.622 1.00 73.49
     ATOM 1872 C GLU 449
    ATOM 1873 O GLU 449
                               49.310 7.973 -1.442 1.00 70.24
     ATOM 1874 N LEU 450
                               51.477 8.113 -0.840 1.00 70.80
                               51.327 7.194 0.285 1.00 68.82
     ATOM 1875 CA LEU 450
    ATOM 1876 CB LEU 450
                               52.693 6.644 0.705 1.00 71.91
45
                               53.428 5.795 -0.336 1.00 76.62
    ATOM 1877 CG LEU 450
    ATOM 1878 CD1 LEU 450
                               54.799 5.414 0.195 1.00 77.95
                               52.617 4.546 -0.662 1.00 76.46
    ATOM 1879 CD2 LEU 450
     ATOM 1880 C LEU 450
                               50.636 7.818 1.492 1.00 66.22
                               50.501 7.181 2.540 1.00 66.01
     ATOM 1881 O LEU 450
50
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	ATOM	1882 N PHE 451	50.189 9.060 1.342 1.00 61.96
	ATOM	1883 CA PHE 451	49.513 9.750 2.428 1.00 58.44
	ATOM	1884 CB PHE 451	50.006 11.204 2.528 1.00 61.34
	ATOM	1885 CG PHE 451	51.466 11.343 2.923 1.00 63.02
5	ATOM	1886 CD1 PHE 451	52.488 10.888 2.077 1.00 62.92
	ATOM	1887 CD2 PHE 451	51.812 11.932 4.146 1.00 63.07
	ATOM	1888 CE1 PHE 451	53.855 11.029 2.437 1.00 65.12
	ATOM	1889 CE2 PHE 451	53.167 12.085 4.531 1.00 64.66
	ATOM	1890 CZ PHE 451	54.195 11.628 3.673 1.00 67.12
10	ATOM	1891 C PHE 451	48.005 9.756 2.219 1.00 56.41
	ATOM	1892 O PHE 451	47.501 10.471 1.350 1.00 56.56
	ATOM	1893 N PRO 452	47.260 8.954 3.009 1.00 53.28
	ATOM	1894 CD PRO 452	47.678 8.027 4.076 1.00 50.46
	ATOM	1895 CA PRO 452	45.797 8.910 2.866 1.00 50.26
15	ATOM	1896 CB PRO 452	45.388 7.976 4.000 1.00 49.19
13	ATOM	1897 CG PRO 452	46.558 7.010 4.039 1.00 45.89
	ATOM	1898 C PRO 452	45.183 10.305 2.974 1.00 49.62
	ATOM	1899 O PRO 452	45.727 11.176 3.644 1.00 52.35
	ATOM	1900 N PRO 453	44.034 10.530 2.313 1.00 51.50
20	ATOM	1900 N TRO 453	43.257 9.585 1.494 1.00 49.66
20	ATOM	1901 CD 1RO 453	43.354 11.830 2.335 1.00 50.89
	ATOM	1902 CA TRO 453	42.101 11.559 1.506 1.00 51.49
	ATOM	1903 CB PRO 453	42.600 10.524 0.521 1.00 50.82
	ATOM	1905 C PRO 453	43.030 12.405 3.706 1.00 50.99
25	ATOM	1906 O PRO 453	43.264 13.588 3.953 1.00 54.17
25	ATOM	1907 N LEU 454	42.479 11.576 4.592 1.00 51.21
	ATOM	1907 N LEU 454	42.112 12.034 5.936 1.00 47.17
	ATOM-	1908 CA LEU 454	41.305 10.951 6.660 1.00 44.44
	ATOM	1910 CG LEU 454	40.748 11.283 8.050 1.00 41.33
30	ATOM	1911 CD1 LEU 454	39.838 12.504 7.978 1.00 35.93
50	ATOM	1912 CD2 LEU 454	39.986 10.072 8.587 1.00 34.79
	ATOM	1912 CD2 EEU 454	43.363 12.380 6.733 1.00 42.25
	ATOM	1914 O LEU 454	43.387 13.357 7.475 1.00 40.82
	ATOM	1915 N PHE 455	44.399 11.567 6.565 1.00 39.29
35	ATOM	1916 CA PHE 455	45.674 11.774 7.240 1.00 41.81
33	ATOM	1917 CB PHE 455	46.655 10.679 6.802 1.00 47.22
	ATOM	1917 CB PHE 455	48.045 10.800 7.407 1.00 56.97
	ATOM	1919 CD1 PHE 455	48.220 10.990 8.785 1.00 57.23
	ATOM	1919 CD1 PHE 455	49.180 10.645 6.597 1.00 59.40
40	ATOM	1921 CE1 PHE 455	49.522 11.030 9.362 1.00 56.58
40	ATOM	1922 CE2 PHE 455	50.487 10.682 7.149 1.00 61.80
	ATOM	1923 CZ PHE 455	50.656 10.870 8.541 1.00 59.94
	ATOM		46,203 13.161 6.892 1.00 45.12
		1924 C PHE 455 1925 O PHE 455	46.558 13.944 7.779 1.00 39.95
45	ATOM	1926 N LEU 456	46.236 13.471 5.592 1.00 43.92
+3	ATOM		46.704 14.767 5.123 1.00 44.08
	ATOM		46.748 14.795 3.593 1.00 50.20
	ATOM		47.796 13.921 2.903 1.00 55.79
	ATOM ATOM	1929 CG LEU 456	47.796 13.921 2.903 1.00 33.79 47.527 13.869 1.408 1.00 54.70
50		1930 CD1 LEU 456	49.187 14.473 3.193 1.00 53.01
50	ATOM	1931 CD2 LEU 456	47.101 17.71 2.173 1.00 33.01

```
45.782 15.871 5.616 1.00 44.65
    ATOM 1932 C LEU 456
                               46.219 16.987 5.887 1.00 45.93
    ATOM 1933 O LEU 456
                               44.500 15.549 5.726 1.00 44.56
    ATOM 1934 N GLU 457
                                43.498 16.504 6.175 1.00 46.37
    ATOM 1935 CA GLU 457
                                42.138 15.854 6.133 1.00 50.16
    ATOM 1936 CB GLU 457
                               43.759 17.039 7.579 1.00 43.60
    ATOM 1937 C GLU 457
    ATOM 1938 O GLU 457
                               43.867 18.245 7.795 1.00 42.69
                               43.847 16.117 8.528 1.00 43.21
    ATOM 1939 N VAL 458
                                44.064 16.446 9.930 1.00 44.98
    ATOM 1940 CA VAL 458
                                44.020 15.159 10.802 1.00 44.83
    ATOM 1941 CB VAL 458
10
                                44.180 15.510 12.277 1.00 49.72
    ATOM 1942 CG1 VAL 458
    ATOM 1943 CG2 VAL 458
                                42,708 14,427 10,567 1,00 40,89
    ATOM 1944 C VAL 458
                               45.368 17.178 10.209 1.00 42.72
                               45.393 18.139 10.974 1.00 42.88
    ATOM 1945 O VAL 458
    ATOM 1946 N PHE 459
                               46,451 16,743 9.574 1.00 44.53
15
    ATOM 1947 CA PHE 459
                                47.741 17.366 9.823 1.00 48.18
    ATOM 1948 CB PHE 459
                                48.784 16.269 10.064 1.00 43.60
    ATOM 1949 CG PHE 459
                                48.374 15.276 11.133 1.00 40.79
                                47.835 14.032 10.783 1.00 41.01
    ATOM 1950 CD1 PHE 459
    ATOM 1951 CD2 PHE 459
                                48.471 15.613 12.492 1.00 39.48
20
                                47.387 13.118 11.776 1.00 40.62
    ATOM 1952 CE1 PHE 459
                                48.032 14.715 13.506 1.00 36.87
    ATOM 1953 CE2 PHE 459
                               47.489 13.463 13.146 1.00 36.39
     ATOM 1954 CZ PHE 459
                               48.234 18.348 8.763 1.00 52.71
     ATOM 1955 C PHE 459
                               49.336 18.878 8.877 1.00 51.34
     ATOM 1956 O PHE 459
25
                               47.397 18.594 7.752 1.00 59.56
     ATOM 1957 N GLU 460
     ATOM 1958 CA GLU 460
                                47.695 19.509 6.647 1.00 66.14
                                47.818 20.944 7.158 1.00 67.76
     ATOM 1959 CB GLU 460
                                46.536 21.511 7.724 1.00 78.99
     ATOM 1960 CG GLU 460
                                46.680 22.965 8.116 1.00 86.08
     ATOM 1961 CD GLU 460
30
                                47.014 23.786 7.237 1.00 87.62
     ATOM 1962 OE1 GLU 460
                                46.460 23.289 9.301 1.00 91.63
     ATOM 1963 OE2 GLU 460
                               48,940 19.163 5.836 1.00 69.17
     ATOM 1964 C GLU 460
                               48.784 18.759 4.660 1.00 69.49
     ATOM 1965 O GLU 460
                                50.057 19.298 6.379 1.00 76.70
     ATOM 1966 OXT GLU 460
35
                              47.283 4.313 16.972 1.00 44.70
     ATOM 1967 C1 TRI
                         1
                              51.052 6.807 13.814 1.00 34.01
     ATOM 1968 C2 TRI
                          1
                              47.289 4.043 15.500 1.00 37.90
     ATOM 1969 C3 TRI
                          1
                              51.936 6.615 12.728 1.00 33.38
     ATOM 1970 C4 TRI
                          1
                              48.462 4.501 14.746 1.00 46.53
40
     ATOM 1971 C5 TRI
                          1
                              52.294 7.653 11.847 1.00 42.90
     ATOM 1972 C6 TRI
                          1
     ATOM 1973 C7 TRI
                          1
                              49.577 5.179 15.334 1.00 34.63
                              51.717 9.015 12.071 1.00 38.34
     ATOM 1974 C8 TRI
                          1
                              49.492 5.383 16.723 1.00 43.89
     ATOM 1975 C9 TRI
                          1
                              50.779 9.237 13.172 1.00 40.43
     ATOM 1976 C10 TRI
                          1
45
                               48.354 4.960 17.533 1.00 41.82
     ATOM 1977 C11 TRI
                          1
                               50,449 8.116 14.055 1.00 35.64
     ATOM 1978 C12 TRI
                          1
                              46.287 3.725 17.959 1.00 36.78
     ATOM 1979 C13 TRI
                          1
     ATOM 1980 C15 TRI
                         1
                              44.825 4.150 17.865 1.00 40.69
                              48.684 4.002 12.609 1.00 40.26
     ATOM 1981 I1 TRI 1
50
```

PCT/US98/25296

	ATOM	1982 I2 TRI I	53.597 7.174 10.336 1.00 46.70	
	ATOM	1983 I3 TRI 1	51.362 6.218 17.644 1.00 36.54	
	ATOM	1984 O3 TRI 1	44.546 5.255 17.329 1.00 54.78	
	ATOM	1985 O2 TRI 1	50.831 5.617 14.667 1.00 28.44	
5	ATOM	1986 O1 TRI 1	52.207 10.160 11.342 1.00 43.65	
3	ATOM	1987 O4 TRI 1	44.021 3.333 18.352 1.00 42.95	
	ATOM	1 AS CAC 501	60.548 16.977 16.916 1.00 65.97	AS
	ATOM	2 AS CAC 502	27.863 16.627 16.796 1.00 89.34	AS
	ATOM	3 AS CAC 503	29.889 28.698 21.811 1.00100.00	AS
10	ATOM	4 AS CAC 504	33.547 24.203 8.880 1.00100.00	AS
10		5 O HOH 505	42.365 8.872 4.597 1.00 53.88	HOH
	ATOM	6 O HOH 506	33.545 30.973 24.585 1.00 40.33	НОН
	ATOM	7 O HOH 507	37.040 1.824 12.671 1.00 61.87	НОН
	ATOM	8 O HOH 508	44.105 4.635 6.023 1.00 40.68	HOH
1.0	ATOM		52.686 13.817 -6.263 1.00 54.00	HOH
15	ATOM		50,186 12.691 -5.997 1.00 55.36	НОН
	ATOM	10 O HOH 510	49.278 18.540 14.006 1.00 34.79	НОН
	ATOM	11 O HOH 511		НОН
	ATOM	12 O HOH 512	25.541 28.885 21.206 1.00 55.42 27.346 31.063 27.398 1.00 58.30	
••	ATOM	13 O HOH 513		HOH
20	ATOM	14 O HOH 514	40.790 19.192 39.234 1.00 50.35	HOH
	ATOM	15 O HOH 515	37.467 0.637 37.293 1.00 37.46	HOH
	ATOM	16 O HOH 516	36.155 3.879 47.189 1.00 61.37	НОН
	ATOM	17 O HOH 517	35.410 5.865 50.995 1.00 63.46	НОН
	ATOM	18 O HOH 518	33.622 5.440 47.570 1.00 53.87	HOH
25	ATOM	19 O HOH 519	64.787 6.888 11.882 1.00 51.15	НОН
	ATOM	20 O HOH 520	61.109 -8.688 27.722 1.00 61.70	НОН
	ATOM	21 O HOH 521	49.869 -5.472 30.343 1.00 40.50	HOH
	ATOM	22 O HOH 522	43.786 -0.987 26.878 1.00 52.16	НОН
	ATOM	23 O HOH 523	41.604 2.361 26.985 1.00 47.90	HOH
30	ATOM	24 O HOH 524	54.405 6.361 39.795 1.00 56.56	НОН
	ATOM	25 O HOH 525	46.088 0.770 33.095 1.00 74.24	HOH
	ATOM	26 O HOH 526	50.481 16.245 15.314 1.00 28.99	HOH
	ATOM	27 O HOH 527	59.788 14.863 21.416 1.00 50.02	HOH
	ATOM	28 O HOH 528	49.282 19.490 32.191 1.00 41.61	НОН
35	ATOM	29 O HOH 529	56.683 10.961 26.733 1.00 34.20	HOH
	ATOM	30 O HOH 530	56.701 9.852 30.561 1.00 51.24	НОН
	ATOM	31 O HOH 531	26.487 13.273 30.591 1.00 43.94	HOH
	ATOM	32 O HOH 532	27.019 25.052 28.330 1.00 54.97	HOH
	ATOM	33 O HOH 533	50.689 1.918 29.551 1.00 30.63	HOH
40	ATOM	34 O HOH 534	47.867 0.200 31.330 1.00 43.14	HOH
	ATOM	35 O HOH 535	61.434 -0.721 23.218 1.00 49.83	HOH
	ATOM	36 O HOH 536	41.969 20.017 20.894 1.00 27.00	HOH
	ATOM	37 O HOH 537	46.897 16.244 15.992 1.00 31.50	HOH
	ATOM	38 O HOH 538	29.796 16.276 27.000 1.00 38.52	HOH
45	<b>ATOM</b>	39 O HOH 539	47.853 23.205 20.217 1.00 44.39	HOH
	<b>ATOM</b>	40 O HOH 540	40.956 24.775 31.717 1.00 50.36	НОН
	ATOM	41 O HOH 541	43.310 1.560 41.912 1.00 43.56	HOH
	END			

PCT/US98/25296

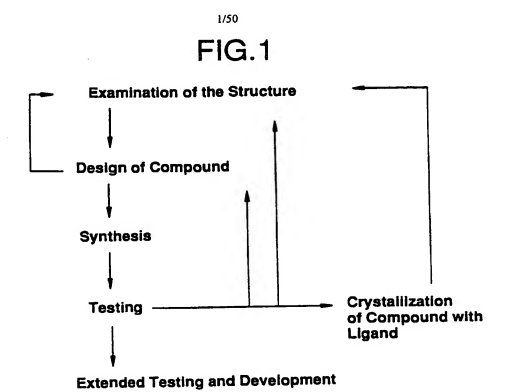


FIG.2

DOMAINS .	NH2-TERMINAL	DNA BINDING	LIGAND BINDING
	: Hypervariable	> 40%	About 20%
FUNCTION:	Transactivation	DNA Binding Dimerization	LIGAND Binding Dimerization Transactivation Nuclear translocation Hsp binding

WO 99/26966

2/50 9 VSCVSGAIPN IPISLDGLLF TSDTLPEVSA SLPEGLDMER RWGQVSQAVE RSSLGPTERT DENNYMEIVN MTELKAKGPR APHVAGGPPS PEVGSPLLCR PAAGPFPGSQ ... METKGYH hGR hPR **HAR** hER hvdr hPPARalpha hPPARbeta hPPARgamma hTRalpha hRARalpha hRARgamma hRXRalpha hRXRbeta rTRalpha hTRbeta

WO 99/26966

3/50 LDSVLDTLLA SATVAESHGL XHDSVRDADY WEFYKTLAG SSVLAQERGD SSPPEKDSGL SLTPGREENP EATRGAGGSS QELLPCLQQD NNRPGILTSD IKTELESKEL SDVEGAYSRA .... MDSKE DEKTODOQSL PRPCQGQDPS NSTOCSSKEK 61 hvdr her hGR hPR P.R. hPPARbeta hPPARgamma hRXRbeta nPPARalpha hTRalpha hRXRalpha hTRbeta hRARalpha hRARgamma

WO 99/26966

4/50 180 SDSGSSVNGG KVGDSSGTAA DFPKGSVSNA . DSKQRRLLV LSPLMSRSGC SYEQQNQQGS HSPAKIYQNV EQLVKFYKGN GHRPSTLSCV NTPL..RSFH **PPAAPATORV** CLFGPELPED PSGPGQSQPS PPACEVTSSW PSLAVASQS. GATVKVSASS 121 hgr hpr hVDR her **LMR** hRXRalpha hPPARbeta hPPARgamma hTRalpha hTRbeta hRARalpha hRARg amma hRXRbeta hPPARalpha rTRalpha

WO 99/26966

SSTIASFGSF PVHSPITQGT ESAGPLLKGK ESIANLNRS. SGETDLKLLE VEEEDSSESE ESPHWSGAPV KPSPQAAAVE FPQQCQISLS VHRAIVK .. S PIHCHEKSPS VCSPLNHTSS VCSPAGINSV ETKVMGNDLG PARQLLLPAS QQPDLSKAVS LSHGLYHGET AHKVLPRGLS 181 hvdr her hGR hPR HAR. rTRalpha hTRalpha hTRbeta hPPARgamma hRASalpha hRARgamma hRXRalpha hRXRbeta hPPARalpha hPPARbeta

5/50

WO 99/26966

300 PVSSPNNVIL **PGRSPLATTV** HLKGQTGING GNVKLYIT. EDSRESAPRV ALVEQDAPHA SPPSHCSVKS THSDVSSEQQ RGSRSHSPAH ASNVGSPLSS PLSSHKSSIS PKSSASTAVS AAPTEKEFPK GGAAACPPGA AAGGVALVPK PLICSPNAEN ... TSVPEN PRALCGAAAG 241 hGR hPR hVDR her HAR HAR hPPARalpha hPPARbeta hPPARgamma hRXRalpha hRXRbeta hTRalpha hRARgamma rTRalpha hTRbeta hRARa I pha

6/50

360 CSPVNNAFSY TASGTSAGSS FAPPRTSPCA .....ET NESPWRSDLL SA.... SPAASTVGSI SGSPGK... GGAGAA.... LNHALLAART RQLLEDESYD NNSRCSVSSP SNTNNRSTLS ..... DQST FDILQDLEFS HDFIHVPILP RSSVSSPANI 301 her hgr hPR hvdr rTRalpha hTRalpha hTRbeta hRARalpha **hRARgamma** hRXRalpha hRXRbeta hPPARalpha hPPARbeta hPPARgamma

7/50

WO 99/26966

8/50 ILPDIKPKIK DNGDLVLSSP SNVILPQVKI SARSPRSYLV TQEKGAQEVP FPKTEEVESA ISNGVIGQLN IVQYIKPEPD GAFSSSCLGG 420 ..... HSW AARPPFLPQR HAEGSVGRWG YSDFQPPALK IKEEEEGAEA GNSNEDCKPL AEPKDDAYPL P. DCAYPPD AGEDDSFLLE IDENCLLSPL TLRDWVPSPD SSTPVAVGDF 361 hgr her hPR hVDR **FAR** hRXRbeta hPPARbeta hPPARgamma hRARgamma hRXRalpha hPPARalpha hTRalpha nRARalpha rTRalpha hTRbeta

FIG.3H

9/50 480 **MSAISVHGVS** TIRVLEVEVD **PASASVSSAS** KDYYSLSGIL YPEGAAYEFN SRSPDSSSPN ......GA VAPYGYTRP. FSTQVNSS .....AA SVPIKQESTK HSCSGTSFKG NPTVNPFPFH DGSYFSFMDD SCILKFPAQD AQVIVMSGQE PLERPLGEVY LDSSKPAVYN NN.... **HDTKHFLPLD** RDGRH..GRD GGGGRRRIIN PGAGARGWIG .....GG GGGEA.... ANIIG EAAVT TVYCQASFPG PLTVNEQLLC EPLNRPQLKI PR. ATPSRPG DTEDLPANNA PGVIKQEKLG AKECIVGSAT ALAGSRSGGG LLHQIQGNEL FPLGPPPPLP TLHTKASGMA EKEDFIELCT AGANPAAFPD NSKINSDSSF 421 hGR hvdr her hPR **h**HR hPPARgamma **hAR** hRARgamma hRXRbeta hPPARbeta hRXRalpha hPPARalpha rTRalpha hTRa1pha hTRbeta hRARa 1 pha

10/50

	481					540
rTRalpha	•		•	•	HEQKPSK	VECGSDPEEN
hTRalpha	•		•	•	HEQKPSK VECGSDPEEN	VECGSDPEEN
hTRbeta	•		MTPNSHTE	NGLTAWDKPK	HCPDREHDWK	LVGHSEACLH
hRARalpha	•		•	•		
hRARgamma	•		•	¥:	ATNKERLFAA	GALGPGSGYP
hRXRalpha	.LISPIGR	LISPIGR., GSMAAPSLHP SLGPGIGSPG .QLHSPISTL SSPINGMGPP FSVISSPHGP	SLGPGIGSPG	.QLHSPISTL	SSPINGHGPP	FSVISSPHGP
hRXRbeta	PLPOGVPP	PLPOGVPP PSPPGPPLPP STAPTLGGSG .APPPP PMPPPPLGSP FPVISSSMGS	STAPTLGGSG	APPPP	PMPPPPLGSP	FPVISSSHGS
hPPARalpha	MVDTESPL	MVDTESPL CPLSPLEAGD LESPLSEEFL QEMGNIQEIS QSIGEDSSGS FGFTEYQYLG	LESPLSEEFL	QEMGNIQEIS	QSIGEDSSGS	FGFTEYQYLG
hPPARbeta	•	HEQPQ EEAP EVREEEEKE EVAEAEGAPE LNGGPQHALP	EEAP	EVREEEEKE	EVAEAEGAPE	LNGGPQHALP
hPPARgamma	•		TEMPFWPINF	GISSVD	LSMADDHSHS	FDIKPFITVD
hVDR	TALSSAGAAE	TALSSAGAAE SGGDEEGSGQ SLEATEEAQL DGPVTTSSTT AVTVEVSAPV VQTVVSKAAI	SLEATEEAQL	DGPVTTSSTT	AUTVEVSAPV	VQTVVSKAAI
her	AAAAANAOVY	AAAAANAQVY GQTGLPYGPG SEAAAFGSNG LGGFPPLNSV SPSPLHLLHP PPQLSPFLQP	SEAAAFGSNG	LCGFPPLNSV	SPSPLMLLHP	PPQLSPFLQP
hgr	TSGGOMYHYD	TSGGQHYHYD HNTASLSQQQ DQ KPIFNVIPP IPVGSEN	DQ	KPIFNVIPP.	IPVGSEN	•
hPR	SSGSTLECIL	SSGSTLECIL YKAEGAPPQQ GPFAPPCKA PGASGCLLPR DGLPSTS	GPFAPPPCKA	PGASGCLLPR	DGLPSTS	•
har	GPPVPGFDGN	GPPVPGFDGN CEGSGFPVGI KQEPDDGSYY PEASIPSSAI VGVNSGGQSF HYRIGAQGTI	KQEPDDGSYY	PEASIPSSAI	<b>VGVNSGGQSF</b>	HYRIGAQGTI
har	•	PQGLAGQE SDFTAPDVWY PGGHVSR VPYPSPT	SDFTAPDVWY	PGGHVSR	VPYPSPT	•

## FIG.3

					11.	/50							
009	SGYI	TEEKKCKGYI	EEIVPSPPSP	EEMVPSSPSP VI.KVPAHPSG	GLHCPPPPGG	YPWVPGSVDE	DOLOHGC. DIS	AGSVGGOGGL	DKGSHAMESA	.SSPPSSSST	DSEASOSPQY	V5551LK5V3	•
	•	OSVSSAQTFQ	AAIETQSSSS	LSVETQSTSS	DVKPPVLGVR	.SSPSSVT	.SSPPSLL	CODITIBLEM	GGRERLASTN	٧	YPPYLNYLRP	YPVLEYIPEN	LF 1D1 16 F
	IX9S	FHLDHDDVND	SNHVASGAGE	QPDLPKEHAS	LPGGGSGPPE			QEYQSAIKVE	TGALACASSA	GYSSPSMRPD	AVLKEGLPQV	GDLSSRRSDG	HRLETARDHV
	LKSSH	LKTSM	Nd	MLSPSFRGLG	GSPQLSS		•	VADYKYDLKL	LTQDGLASLA	NEPCRTUFSN	NGLPQLGYQA	NTLVESWKSH	PYGD
	RKN. GQCP	RKN.GQCS	LAN. EQSSEN	ALR.GSPPFE	. TTPTLGFST GSPQLSS FAMFVSSSE LIGHTERS GLHCPPPFGG	TDTLSPA	LSRS	DIPFTRADPH	PITVQACPQV	HGQQVPYYLE NEPSGYIVKE AGFFnf introduced V SSPPSSSST	. ASAAAAGA APALYPALGI NGIPQIGYQA AVIKEGIPQV YPPYINYIRP DSEASQSPQY	SLSRSARDQS FQHLSSFPPV NTLVESWKSH GDLSSRRSDG YPVLEYIPEN VSSSILKSVS	CVKSEMGP WMDSYSGPYGD MRLETAKUHV LFIDIIFF
	SARSPOGKRK RKN.GQCPLKSSH	SARSPOGKRK RKN.GQCSLKTSM	RKSHSEKKSI LINI ELSSEN LIKITING SANDETQSSS EEIVPSPPSP	GAGFPFAFPG ALR. GSPPFE HLSPSFRGLG QPDLPKEHAS LSVETQSTSS EEHVPSSPSP		SCPGSDGSVI TDTLSPA	SSSYTDLSRS	FSSISAPHYE DIPETRADPH VADYKYDLKL QEYQSAIKVE PASPFIISEN AKLINKING	SVSPAQQISV PITVQACPQV LTQDGLASLH TGRLAXYSSL CELLISTIN DKGSHAMESA	HCQQVPYYLE	. ASAAAAGA	SLSRSARDQS	CVKSEMGP
	rTRalpha	hTRalpha	hTRbeta	nkaka i pina hRARgamma	hRXRalpha	hRXRbeta	hPPARbeta	PPARgamma	hvdr	her	hgr RPA	P.M.R.	hAR

12/50 099 QAVLQPQMSA DL..... DK.... TN.... DR. DL.. TYSCRD NK. NH..VYTCHR KLVYD...KC KLIYD...RC HN..DYHCPA QHNYLCAGRN KOKYLCASRN NLHPTYSCKY NLHPTYSCKY NH.. VYTCHR DL..TYTCRD KLEYE...KC SVQTQLQAPA QHNYLCAGRN QHNYLCAGRN NLHPSYSCKY KGFFRRTIOK KGFFRRSIQK KGFFRRTIQK KGFFRRTIQK KAFFKRSIQG HYGALTCGSC KVFFKRAAEG KGFFRRSIQK KGFFKRTVRK KGFFKRTIRK KGFFRATIRL KGFFRRTIRM KGFFRRTIRL AGLQAATVLN KVFFKRAVEG KVFFKRAMEG KVFFKRAVEG HYRCITCEGC HYGVSSCEGC HYGVWSCEGC HYGVLTCGSC HYGWYCGSC HY RCITCEGC HYRCITCEGC HYGVSACEGC HYGVYSCEGC HYGVYSCEGC HYGVHACEGC HYGVHACEGC HYGVLTCGSC PAGGLLKLPF HYGVHACEGC WCGDKATGY **FVCNDKSSGY** AICGDRSSGK LICCDEASGC LVCGDEASGC LICGDKASGC VVCGDKATGY RICGDKASGY RVCGDKASGF LVCSDEASGC WCGDKATGY AICGDRSSGK RVCGDKASGF ATLPGLAAAS AVCNDYASGY FVCQDKSSGY ATTGPPPKLC IGSSRPSKIC NHASFTKHIC AVLTLPTATV SFESTPOKIC ....PQKTC PSYLDKDELC PPPPRVYKPC PGAG. . KRLC SPSGALNIEC KET...RYC PSYLDKDEQC PSYLDKDEQC PPLPRIYKPC ASCGSLNMEC PSNSLMAIEC 601 hvdr her hGR hPR hRXRalpha hPPARgamma FAH RAH rTRalpha hTRalpha hTRbeta hRARalpha hRARgamma hRXRbeta hPPARalpha hPPARbeta

FIG.3K

13/50

<b>L</b>	<b>~</b>	¥	ρţ	Ē	Æ		ري پ	z	လှ	۲	ŝ	.•	>	Ø	S.	
RRKEEHI	RRK. EEHI	RRR EELQ	KKKEVPK	KKKEVKE	EVESTSS	DGECAGG	LKA. EILT	LVAGLTA	LLAEI.S	<b>QPQFISSLT</b>	EGRGEVG	GIQQATT	WWRALDA	GIHEEQP	LQEEGEA	
RKLIEQNRER	RKLIEQNRER	RKL I EENREK	RNK	RNK KKKEVKEE	KDRNEN EVESTSSA	KDK. DG DGE CAGGA	.RHPRSEKAK LKAEILTC	.RHPEAEKRK LVAGLTAN	. RHPQAEKEK LLA EI.SS	AIISAASLGA	LKHKRQRDDG	RKTKKKIK GIQQAIT.	RKFKKFNKVR VVRALDAV	RKSKKLGKLK GIH EEQPQ	RKLKKLGNLK LQEEGEAS	
VLDDSKRVAK	VLDDSKRVAK	VLDDSKRLAK	VRND	VRND	VQEERQRG	VQEERQRG	IRFG	IRFG	IRFG	LQTAGLSINP	IRKDRRGGRH	•		•	•	
CIAVGHAMDL	CIAVGHAHDL	CIYVGHATDL	CFEVGHSKES	CFEVGMSKEA	CLAHGHKREA	CLATGHKREA	CLSVGMSHNA	CLALGMSHNA	CLAVGHSHNA	ASEPSVSVAT	CYEVGHMKGG	CLQAGHNLEA	CCQAGHVLGG	CLQAGMNLGA	CYEAGHTLGA	
.CCVIDKITR NOCOLCRFKK CIAVGHAHDL VLDDSKRVAK RKLIEQNRER RRKEEHIR	.CCVIDKITR NQCQLCRFKK CIAVGMAHDL VLDDSKRVAK RKLIEQNRER RRKEEHIR	KCVIDKVTR NOCOECRFKK CIYVGHATDL VLDDSKRLAK RKLIEENREK RRREELQK	NCIINKVIR NRCQYCRLQK CFEVGHSKES VRNDRNK KKKEVPKP	.NCIINKVIR NRCQYCRLQK CFEVGMSKEA VRND	DCLIDKROR NRCQYCRYOK CLAHGHKREA VQEERQRG.	.DCTVDKRQR NRCQYCRYQK CLATGMKREA VQEERQRG.	SCKIQKKNR NKCQYCRFHK CLSVGMSHNA IRFG	SCKIQKKNR NKCQYCRFQK CLALGMSHNA IRFG	NCRIHKKSR NKCQYCRFQK CLAVGMSHNA IRFG	loamootott aattasivok asepsvsvat lotaglsinp aiisaaslga opofisslit	. QCTIDKNRR KSCQACRLRK CYEVGHHKGG IRKDRRGGRH LKHKRQRDDG EGRGEVGS	.CIIDKIRR KNCPACRYRK CLOAGHNLEA	.CIVDKIRR KNCPACRLRK CCQAGHVLGG	CIIDKIRR KNCPACRLQK CLQAGHNLGA	CTIDKFRR KNCPSCRLRK CYEAGHTLGA	•
. CCVIDKITR	.CCVIDKITR	.KCVIDKVTR	.NCIINKVTR	.NCIINKVTR	. DCLIDKROR	.DCTVDKRQR	. SCKIQKKNR	. SCKIQKKNR	. NCRIHKKSR	LOAMOOTOTT	.QCTIDKNRR	CIIDKIRR	CIVDKIRR	CIIDKIRR	CTIDKFRR	
rTRalpha	hTRalpha	hTRbeta	hRARalpha	hRARgamma	hRXRalpha	hRXRbeta	hPPARalpha	hPPARbeta	hPPARgamma	hvdr	her	hGR	hPR	hmr	har	

780

PDDIGOSPIV

SHWKORRKFL

EAHRSTNAOG

EEWDLIHVAT EEWDLIHIAT

SLOCIRPEPTP SLOGRPEPTP

rTRalpha hTRalpha hTRbeta hRARalpha hRARgamma hRXRalpha hRXRbeta hPPARalpha hPPARbeta

minimal start site 725

PEDIGOAPIV TNNSSEQRV.

SHWKQRRKFL PDDICQSPIV KNSLALSLTA DOMVSALLDA EPPILYSE.. CQL...GKYT TNSSADHRV. CQL...GKYT SHWKOKPKFL GLNPS.... TKAKARAILT PLLVNPASLA PTLVS.... GGSGS.... NKVKARVILS TKKKARSILT TALVPQLSTI FSPGQDIQLI PPLIN.... VSHIEGYECQ PIFLN.... KAHQETFPAL KAHQETFPSL ETYVE..ANH DOCVEGPGGT EAYLKNFN.H NAYLKNFN. H DSYIKSFP.L NAQCQVIGTL IVPATLPQLT IAPAKEPSVN EAHRSTNAQG EAHVATNAOG

PSPLMIKRSK

AGDHRAANLW

TSAMSN

TPI

**hVDR** her hGR hPR **HAR** 

ALPOPLGVPN ddddddd000

ETSENPGNKT ESQALSQRFT POSPEEGITY

..GVSQ

14/50

IHDMETICHA

GKASNNPPFV

....SP..

QLEELITKVS

EAELAVEPKT EAELAVEOKS DLKSLAKRIY DLKAFSKHIY DLRALAKHLY VAGLTSQLIT

EVGELIEKVR

EEWELIKTVT

SIGHKPEPTD

ECSESYTLTP GSPUSYELSP NEDMPVERIL PEEMPVDRIL EHD LEDSETA IHDIETLWOA IYDHNSLHHG

GKASHTAPFV

GKTTDKSPFV GAAAASA... LL.... LL

• • • • • • • • • •

SRALTPSPVM

TEETTOKLT

EGSQYNPQVA

DIDOLNPESA

hPPARgamma

15/50

## FIG.3N

16/50

LKNGGLGV VSDAIFELGK	N GGLGV VSDAIFELGK	NGGLGV VSDAIFDLGM	.N. AGFGP LIDLVFAFAN	NAGFGP LTDLVFAFAG	VGAI FDRVLTELVS	SAI FDRVLTELVS	SLRKPFCD IMEPKFDFAM	SLRKPFSD IIEPKFEFAV	SLRKPFGD FMEPKFEFAV	NLEEIREFAK	FDMLLAT. SS	LPCHYD QCKHMLYVSS	ESSFYS LCLTHWQIPQ	QSAMYE LCQGMHQISL	S QCVRMRHLSQ	
LKNGGLGV	NGGLGV	NGGLGV	1 AGFGP	AGEGP	VGAI	GAI	ဥ	Ω	$\sim$					4.5	t/O	
LK.			-	. N .	S.AG	S.AGV	SLRKPF	SLRKPFS	SLRKPFG	PSAVKDEEAI		LPCHYD	ESSFYS	QSAHYE	KSRMY	
MTVKRKQ	HAVKREQLK	MAVIRGOLK.	LTLNRTQMH.	LTLNRTQMH.	LHVHRNSAH.	LHVHRNSAH.	F. ITREFLK.	F. VTREFLR.	F.HTREFLK.	HPTVGQLVNK	LLLDRNQGK.	LIINEQRHT.	LILNEQRMK.	LVFNEEKHH.	LVFNEYRHH.	
ESDTLTLSGE	ESDTLTLSGE	ESETLTLNGE	EQDIMIFSDG	EQDIMIFSDG	VKDGILLATG	VRDGILLATG	DCHLVAYGNG	DGLLVANGSG	DGVLISEGQG	TAAGVIACGE	HPGKLLFAPN	SANLLCFAPD	SCOMLYFAPD	NSQFLYFAPD	NSRMLYFAPD	
RAAVRYDP	RAAVRYDP	RAAVRYDP	RICTRYTP	RICTRYTP	SFSHRSIA	SFSHRSID	HLSSVMNK	MLASIVNK	HLASLMNK	PSVVKPVTSL	GLVWRSHE	ALGWRSYROS	GLGWRSYKHV	ALSWRSYKHT	AMGWRSFINV	
KGCCHEIMSL	KGCCMEIMSL	KGCCMEIMSL	KAACLDILIL	KAACLDILML	RAGWNELLIA	RAGWNELLIA	KYGUYEAIFA	KYGVHEAIFA	KYGVHEIIYT	APSKVIIAPO	ECAMLEILMI	OXSHMFLMAF	OYSWMSTHVF	OYSWMCLSSF	OY SWHGLHVF	t
rTRalpha	hTRalpha	hTRheta	hRARaloha	hRARGamma.	hRXRalpha	hRXRheta	hppagalnha	hpparheta	TOD BUTTER	HVDR	H H H	854	hPR	A A A	har har	
			KGCCHEIHSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIHSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIHSL RAAVRYDP ESETLTLNGE HAVIRGQLK.	KGCCHEIHSL RAAVRYDP ESDTLTLSGE HTVKRKQLK. KGCCHEIHSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIHSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQHH.	KGCCHEIHSL RAAVRYDP ESDTLTLSGE HTVKRKQLK. KGCCHEIHSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIHSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQMH.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE MAVKREQLK. KGCCHEIMSL RAAVRYDP ESETLTLNGE MAVIRGQLK. KAACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQMH. KAACLDILML RICTRYTP EQDTHTFSDG LTLNRTQMH. RAGWNELLIA SFSHRSIA VKDGILLATG LHVHRNSAH.	KGCCHEIHSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIHSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIHSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQHH. KAACLDILMA RICTRYTP EQDTHTFSDG LTLNRTQHH. RAGWNELLIA SFSHRSIA VKDGILLATG LHVHRNSAH.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIMSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQMH. KAACLDILIAL RICTRYTP EQDTHTFSDG LTLNRTQMH. RAGWNELLIA SFSHRSIA VKDGILLATG LHVHRNSAH. RAGWNELLIA SFSHRSID VRDGILLATG LHVHRNSAH. KYGVYEAIFA HLSSVHNK DGHLVAXGNG F.ITREFLK.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE MAVKREQLK. KGACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQHH. KAACLDILML RICTRYTP EQDTHTFSDG LTLNRTQHH. RAGWNELLIA SFSHRSIP VKDGILLATG LHVHRNSAH. RAGWNELLIA SFSHRSID VKDGILLATG LHVHRNSAH. KYGVYEAIFA HLSSVHNK DGHLVANGNG F.ITREFLK.	KGCCHEIHSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIHSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIHSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILLL RICTRYTP EQDTHTFSDG LTLNRTQMH. RAACLDILLA SFSHRSIP EQDTHTFSDG LTLNRTQMH. RAGWNELLIA SFSHRSID VKDGILLATG LHVHRNSAH. RYGVYEAIFA HLSSVMNK DGHLVAXGNG F.ITREFLK. KYGVYEAIFA MLASIVNK DGLLVANGSG F.YTREFLR.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE MAVKREQLK. KGACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQHH. KAACLDILML RICTRYTP EQDTHTFSDG LTLNRTQHH. RAGWNELLIA SFSHRSIP VKDGILLATG LHVHRNSAH. RAGWNELLIA SFSHRSID VKDGILLATG LHVHRNSAH. KYGVYEAIFA HLSSVHNK DGHLVANGNG F.ITREFLK. KYGVHEAIFY HLASIVNK DGLLVANGSG F.WTREFLR. KYGVHEIIYT MLASIMNK DGVLISEGQG F.WTREFLK.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQHH. KAACLDILIL SFSHRSIP EQDTHTFSDG LTLNRTQHH. RAGWNELLIA SFSHRSIP VKDGILLATG LHVHRNSAH. RAGWNELLIA SFSHRSID VKDGILLATG LHVHRNSAH. KYGVYEAIFA HLSSVHNK DGHLVAXGNG F.ITREFLK. KYGVHEIIYT MLASIWNK DGLLVANGSG F.VTREFLR. KYGVHEIIYT MLASLMNK DGVLISEGQG F.HTREFLK. APSKVIIAPQ PSVVKPVTSL TAAGVIACGE MPTVGQLVNK ECAWLEILMI GLVWRSHE HPGKLLFAPN LLLDRNQGK.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE MAVKREQLK. KGCCHEIMSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILLL RICTRYTP EQDTHTFSDG LTLNRTQMH. RAACLDILLA SFSHRSIP EQDTHTFSDG LTLNRTQMH. RAGWNELLIA SFSHRSIP VXDGILLATG LHVHRNSAH. RXGVYEAIFA HLSSVHNK DGHLVAXGNG F.ITREFLK. KYGVHEAIFA HLSSVHNK DGLLVANGSG F.UTREFLK. KYGVHEIIYT HLASIWNK DGVLISEGQG F.HTREFLK. APSKVIIAPQ PSVVKPVTSL TAAGVIACGE MPTVCQLVNK ECAWLEILHI GLVWRSHE HPGKLLFAPN LLLDRNQGK.	KGCCHEIMSL RAAVRYDP ESDTLTLSGE MTVKRKQLK. KGCCHEIMSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGCCHEIMSL RAAVRYDP ESETLTLNGE HAVIRGQLK. KAACLDILLL RICTRYTP EQDTHTFSDG LTLNRTQHH. RAACLDILLAL SFSHRSIP VKDGILLATG LHVHRNSAH. RAGWNELLIA SFSHRSID VKDGILLATG LHVHRNSAH. RYGVYEAIFA HLSSVHNK DGHLVAXGNG F.ITREFLK. KYGVYEAIFA MLASIVNK DGLLVANGSG F.VTREFLK. KYGVHEIIYT MLASLMNK DGVLISEGQG F.HTREFLK. APSKVIIAPQ PSVVKPVTSL TAAGVIACGE HPTVGQLVNK ECAWLEILHI GLVWRSHE HPGKLLFAPN LLLDRNQGK. QYSWHFLMAF ALGWRSYRQS SANLLCFAPD LIINEQRMT. OYSWMSLHVF GLGWRSYKHV SGQHLYFAPD LILNEQRMK.		KGCCHEIHSL RAAVRYDP ESDTLTLSGE HTVKRKQLK. KGCCHEIHSL RAAVRYDP ESDTLTLSGE HAVKREQLK. KGACLDILIL RICTRYTP EQDTHTFSDG LTLNRTQHH. KAACLDILLIL SFSHRSIP EQDTHTFSDG LTLNRTQHH. RAGWNELLIA SFSHRSIP VKDGILLATG LHVHRNSAH. RAGWNELLIA SFSHRSID VKDGILLATG LHVHRNSAH. RYGVYEAIFA HLSSVHNK DGHLVAXGNG F.ITREFLK. KYGVHEIIYT HLSSLMNK DGLLVANGSG F.WTREFLK. KYGVHEIIYT HLSSLMNK DGLLVANGSG F.WTREFLK. APSKVIIAPQ PSVVKPVTSL TAAGVIACGE HPTVGQLVNK ECAWLEILHI GLVWRSHE HPGKLLFAPN LLLDRNQGK. QYSWHFLHAF ALGWRSYRUS SANLLCFAPD LIINEQRHT. QYSWHCLSSF ALSWRSYKHT NSQFLYFAPD LVFNEYRHH.

FIG.30

960

LA...FEHYV LA...FEHYV LA...FEHYI EA...LKVYV

..RSGLLCVD KIEKSQEAYL ..RSGLLCVD KIEKSQEAYL

MSTD. MSTD. MSSD.

EVALLQAVLL EVALLQAVLL EVALLQAVLL ETGILSAICL ETGLLSAICL ETGLLSAICL

SLSAFNLDDT

rTRalpha hTRalpha

901

SLSAFNLDDT

RIEKYQDSFL RVDMLQEPLL KVDKLQEPLL EVEALREKVY EVEVLREKVY

.. RPGLACVE

.. RQDLEQPD

ICGD.....
FNPDS....
CCGD....

QLLPLEMDDA QLLPLEMDDT

hRARa 1 pha

hTRbeta

...KGLSNPA

..RPGLMNVP RVEAIQDTIL
..RPGLLNVK PIEDIQDNLL
SAICREEKLD ITPKSAQKLK
STLKSLEEKD HIHRVLDKIT
VPKDGLKSQE LFDEIRMTYI
IPLEGLRSQT QFEEMRSSYI
IPKDGLKSQA AFEEMRTNYI
IPVDGLKNQK FFDELRMNYI

LNT.....

FSI

KELRKHVTKC KELDRI I ACK

DTLIHLMAKA KELGKAIVKR

FIG.3P

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17/50

AS...LEAYC

AS...LETYC HV...LRLHL RA...LEFHL QA...LELQL PVLERHLAEA

HIEKMQEGIV

.. RPGLLNVG

ELGCLRAIIL

KHRDMQMDKT KHRDMRMDKT

hRXRalpha hRXRbeta hPPARalpha

hRARgamma

DISLEVAAII

KFNALELDDS

LSGD..... TATECPAYSQ LNSGVYTFLS LSS.....

KFNALELDDS

KFNALELDDS

hPPARbeta

hPPARgamma

NFKIRRLSLG

hvor her hor hpr

DLALFIAAII DLAIFIAVII LTQTQVGQAL EFUCLKSIIL EYLCHKTLLL EFLCHKVLLL EYTIHKVLLL EFLCHKALLL

RFRHHNLOGE

ELHRLQVSYE

**QFVRLQLTFE** 

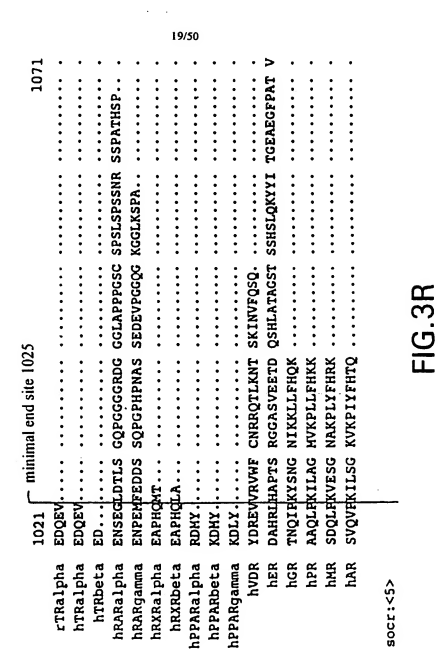
**EFGWLQITPQ** 

**EFVKLQVSQE** 

EFPAMLVEII EITEIAKELN **EFPEHMSEVI** DFPEMMAEII CHASRFL..H MKVEC..PTE LFPPLFLEVF LFPPLFLEVF LLPPLFLEVF SCHPLLQEIY PLYDLLLEML EFPEMLAEII M. PPLIREML SLHPLLQEIY M. PPLIQEML PIDTFLHEHL PIDTFLHEM ALHPLLQEIY IKKTE..SDA IQSRA..LSV RESHA..LKV IKSHM..VSV KGAERVI..T LKMEI..PGS FKL. I..GDT KCLEHLF..F FKL.I..GDT EHAQHMQ..R IKKTE..TET EHVQLLH..V IKKTE..TDM FEKNSLPTGQ CHASRFL..H MKVEC..PTE CHASRFL..H MKVEC..PTE LKMEI..PGP MKC. K.. NVV LD.KT..HSI RKNPTSCSRR FYQLT.... K LLDSVQPIAR ELHQFTFDLL KITDLRGIST KGAERAI..T EHAQLVQ..I ILSHIRHMSN KGMEHLY..S NLLNYCFOTF **QLHLYCLNTF** DLLEFCFYTF KCLEHLF..F POAIEVLNTY LLDSMHEVVE FYQLT...K LLDSMHDLVS RLPALRSIGL RLPALRSIGL KMADLRQLVT KHTDLRQIVT LMEFVGGEPS KKRKRRTSFT QKGVVSSSQR FYQLT...K LLDNLHDLVK KVTDLRMIGA KVTDLRMIGA KITDLRSISA KMADLRQLVT KVTDLRMIGA FYQLT...K WPKLL...H WPKLL...H FPKML...H FPKLL...Q FAKVL...Q LAQLL...L WPKLL...H FPRHL...H FAKLL...L FAKLL...L FPKLL...Q QANHPDAQYL EGNSSQNWQR PNNSGQSWQR RKRRPSRPHH KHKYPEQPGR KOKYPEQQGR QSNHPDDIFL KLNHPESSQL RRRPSQPYH ELWNQKGQQN GLTLQQQHQR NHRKHNI PHF NHRKHNIPHF NYRKHHVTHF 961 hvdr her hGR hPR **h**HR hRXRbeta hPPARalpha hPPARbeta hPPARgamma rTRalpha hTRalpha hTRbeta hRARa 1pha **hRARgamma** hRXRalpha

18/50

PCT/US98/25296



PCT/US98/25296

20/50

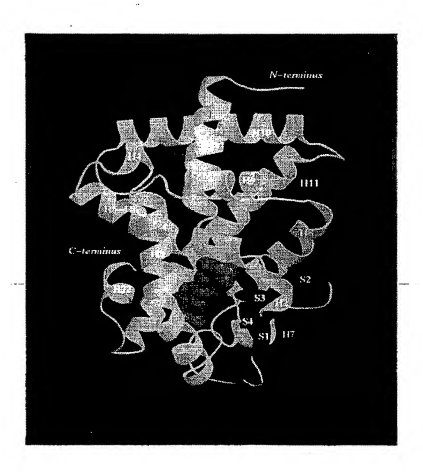
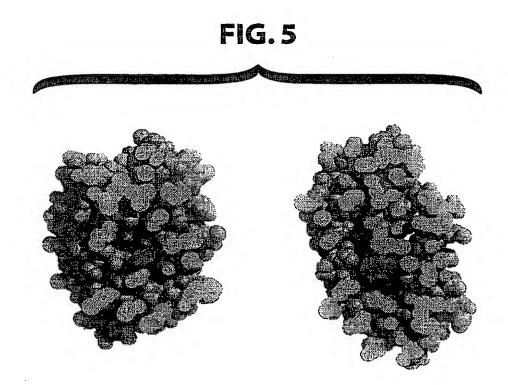


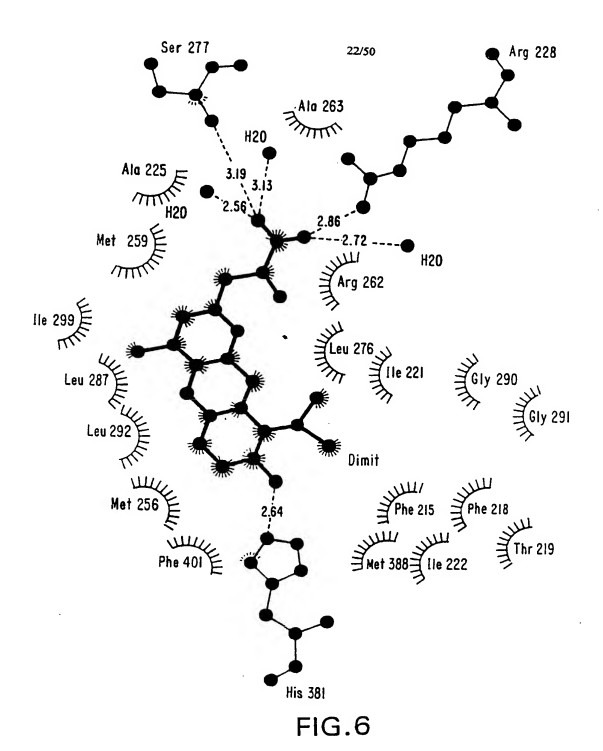
FIG. 4

PCT/US98/25296

21/50



PCT/US98/25296



WO 99/26966

23/50



FIG. 7

PCT/US98/25296

24/50

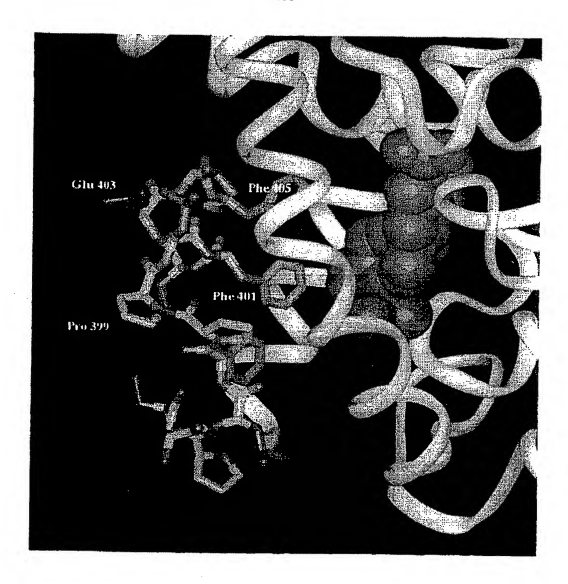


FIG.8

WO 99/26966 PCT/US98/25296

25/50

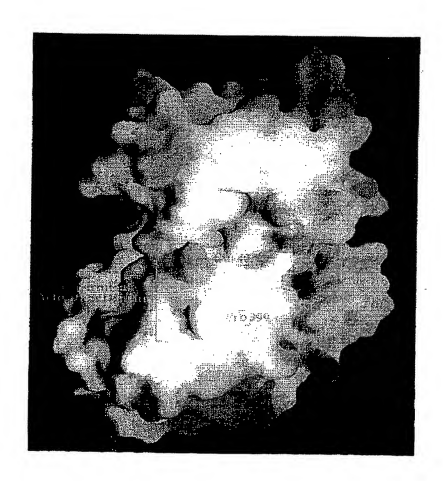


FIG. 9

PCT/US98/25296

## **AGONISTS**

26/50

<u>ANTAGONISTS</u>

Retinoic Acid

HO Estradiol

HO

N(CH<sub>3</sub>)<sub>2</sub>

Diethylstilbestrol

Progesterone

(CH<sub>3</sub>)<sub>2</sub>N OH OH RU 486

FIG.10

shows position of extension group

PCT/US98/25296

Compound	RCOX
TSI	Ph <sub>2</sub> CHCO <sub>2</sub> NHS
TS2	C <sub>16</sub> H <sub>33</sub> CO <sub>2</sub> NHS
TS3 ·	FMOC-CI
TS4	tB <b>0C</b> <sub>2</sub> 0
TS5	tB <b>0</b> C20

## FIG.11

PCT/US98/25296

PCT/US98/25296

29/50

30/50

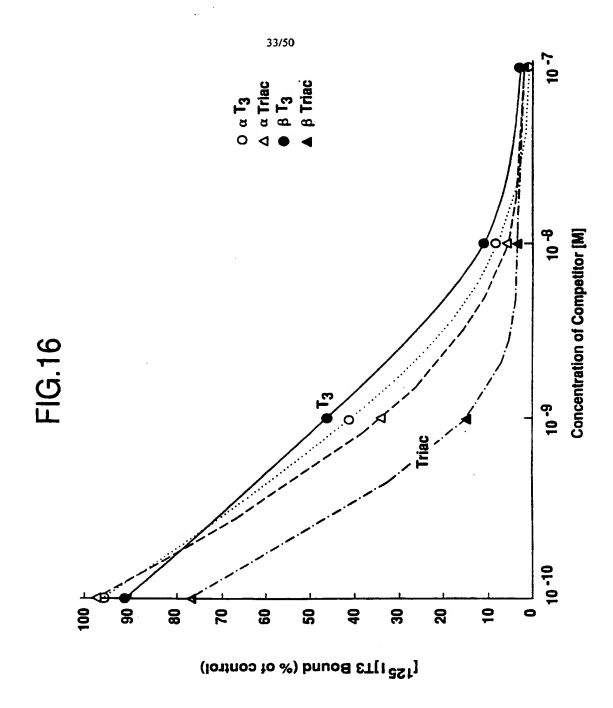
31/50

PCT/US98/25296

$$\begin{array}{c} & & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

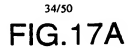
FIG. 15

PCT/US98/25296



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WO 99/26966



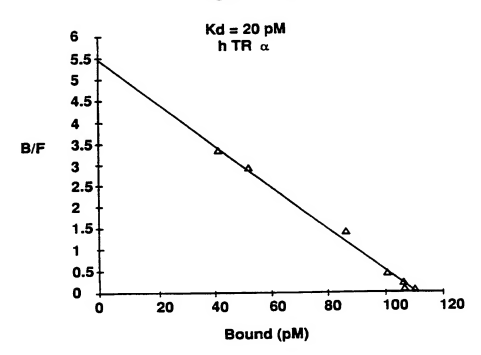
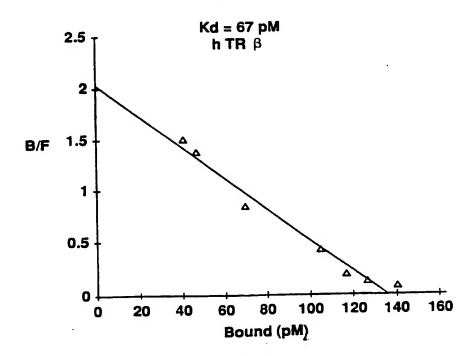
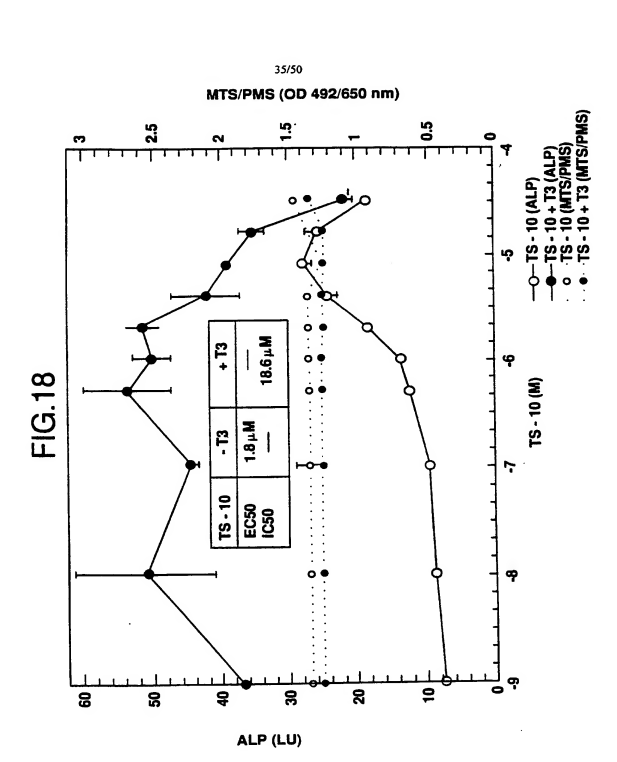


FIG.17B

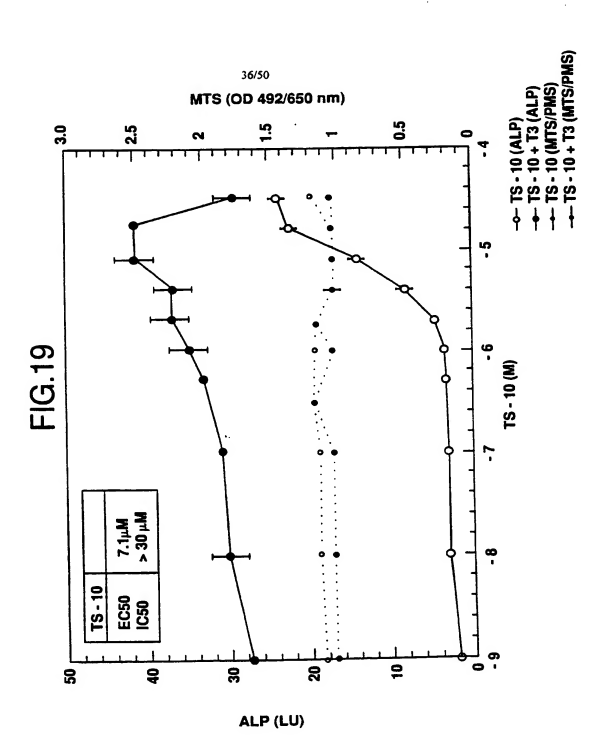


SUBSTITUTE SHEET (RULE 26)

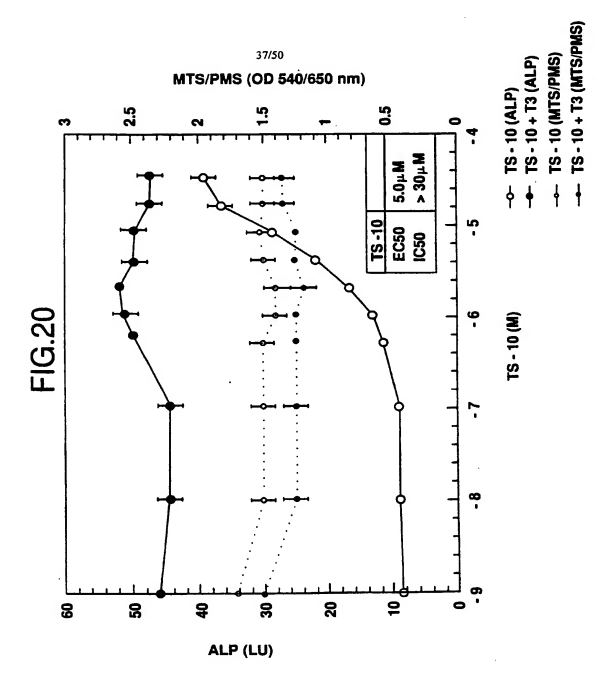
WO 99/26966



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SUBSTITUTE SHEET (RULE 26)



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PCT/US98/25296

38/50

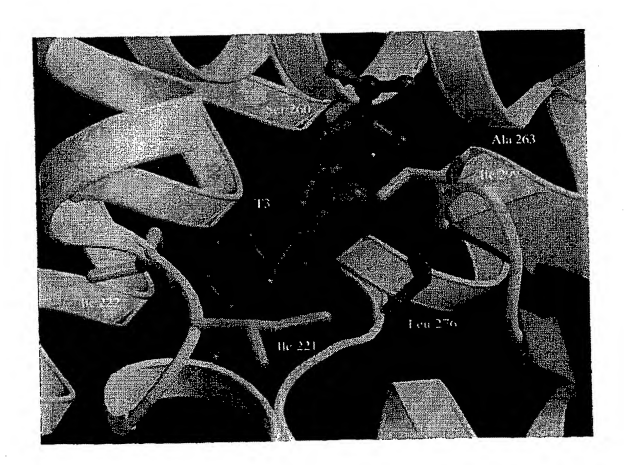


FIG. 21

39/50

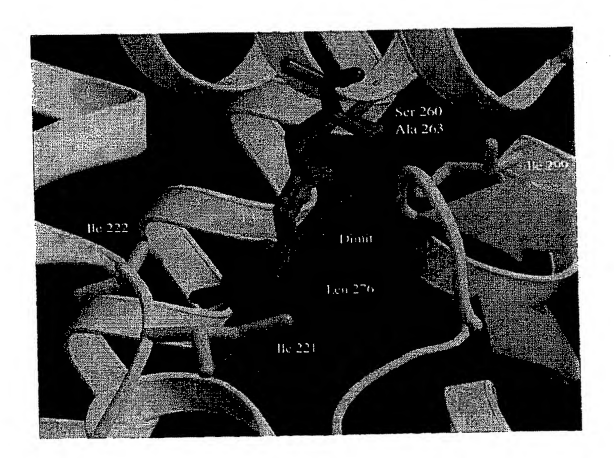


FIG. 22

40/50

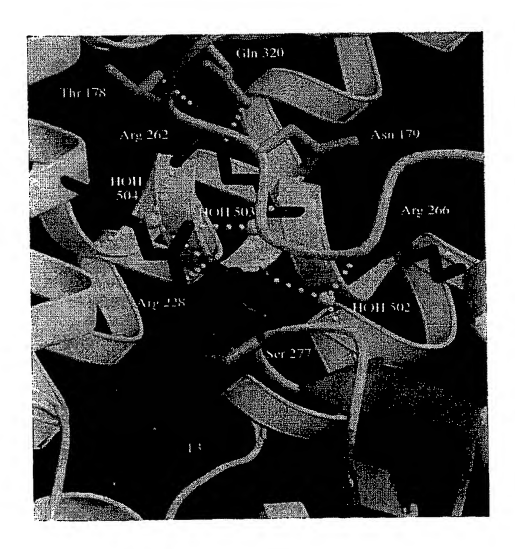


FIG. 23

PCT/US98/25296

41/50

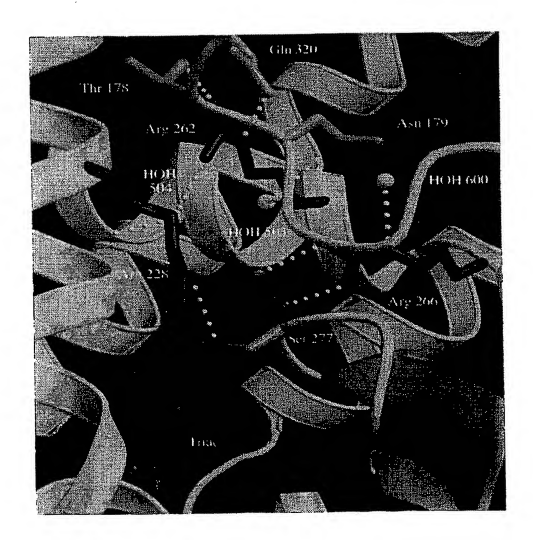


FIG. 24

42/50

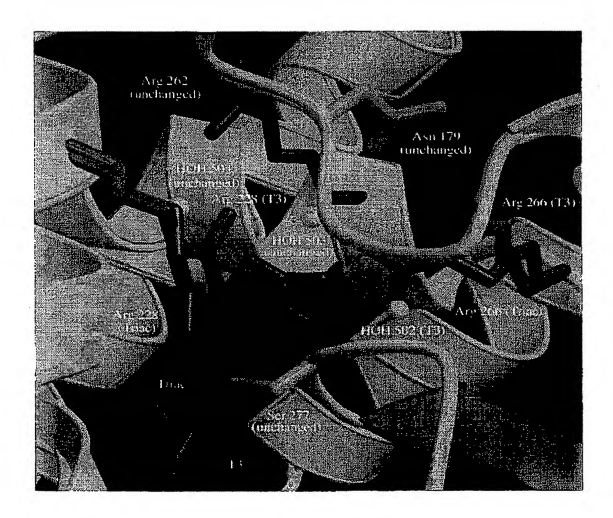
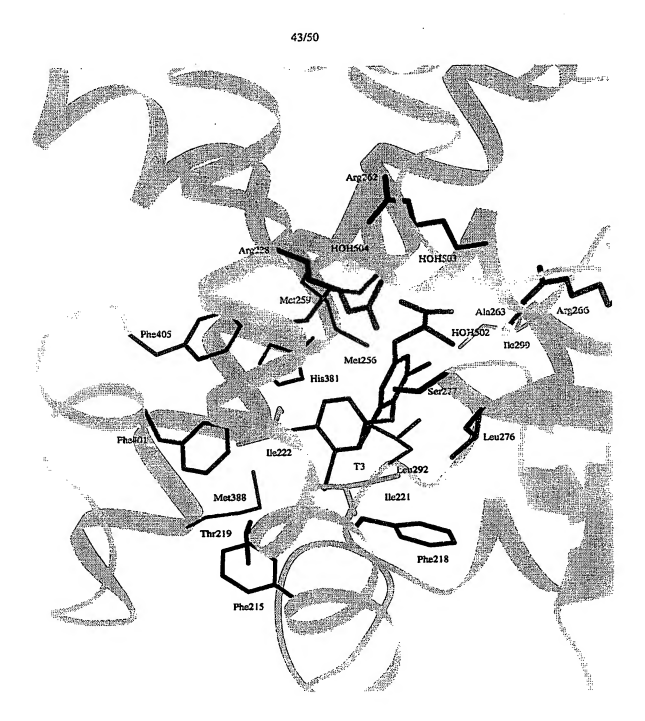
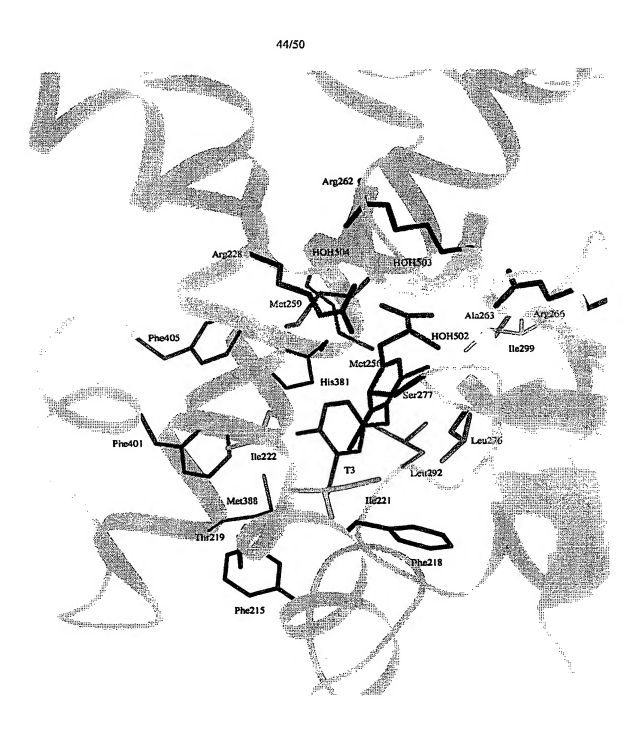


FIG. 25



**FIG. 26A** 



**FIG. 26B** 

PCT/US98/25296

45/50

Thyroid Hormone Receptor Beta with GC1

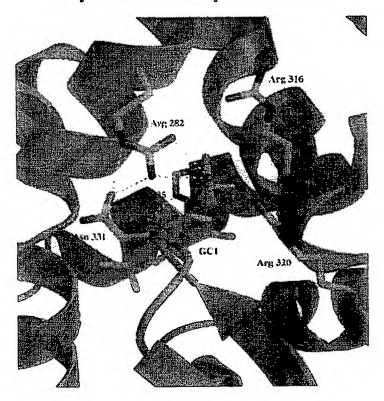


FIG. 27

46/50

Thyroid Hormone Receptor Beta with Triac

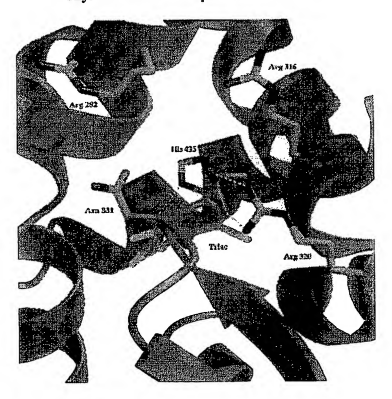


FIG. 28

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PCT/US98/25296

47/50

## Structural Differences Between TR-b with GC1 and TR-a with Dimit

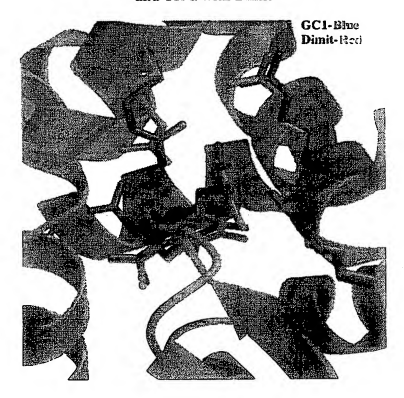


FIG. 29

PCT/US98/25296

48/50

#### Structural Differences between TR LBD isoforms with Triac

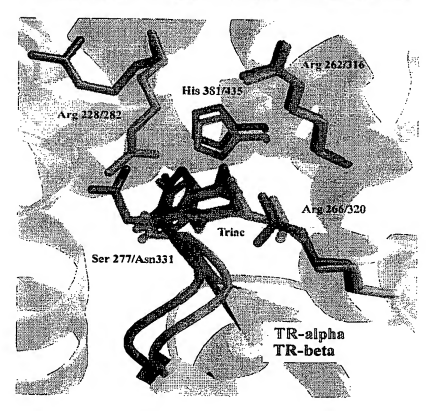
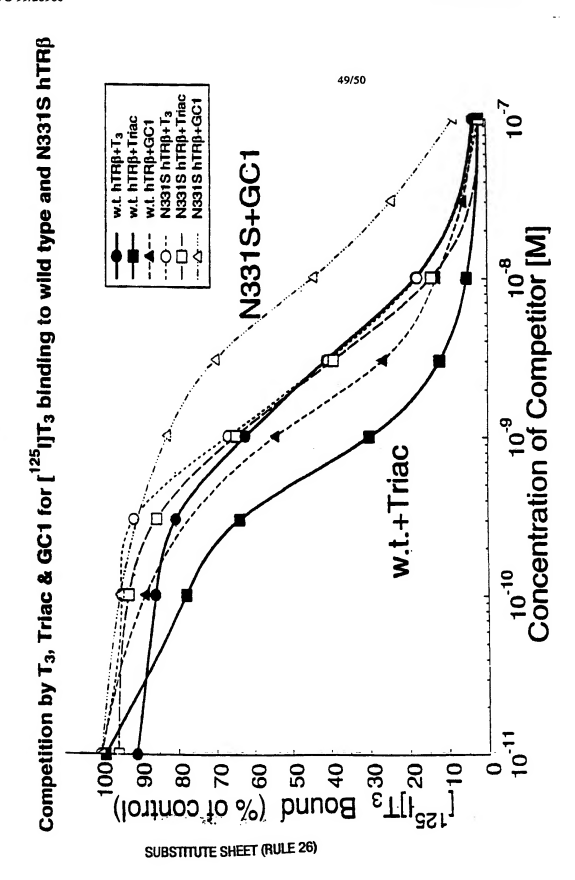


FIG. 30

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50/50
Atomic Numbering for Thyronine-like Ligands

$$R5$$
 $C10 - C12$ 
 $C9 = C11$ 
 $R4' - C8$ 
 $C2 - X - C7$ 
 $C1 - R1$ 
 $C6 = C4$ 
 $C5 - C3$ 
 $C3$ 

Ligand	R1	R3	R5	X	R3'	R4'
Dimit	amino propionic	C19	C20	02	iPr	01
IpBr,	amino propionic	BR1	BR2	02	iPr	01
Ť.	amino propionic	I1	13	02	12	<b>O</b> 1
Triac	acetic acid	<b>I</b> 1	13	02	12	<b>O</b> 1
GC1	oxyacetic acid	C19	C20	C21	iPr	<b>O</b> 1

**FIG.32** 

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